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Human Nucleic Acid Sequences from Hysteromyomic Tissue

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The invention relates to human nucleic acid sequences from hysteromyomic tissue, which code for gene products or portions thereof, their functional genes that code at least one bioactive polypeptide and their use.

In addition, the invention relates to the polypeptides that can be obtained by way of the sequences and their use.

One of the main causes of death by cancer in women is hysteromyoma, for control of which new therapies are necessary. Previously used therapies, such as, e.g., chemotherapy, hormone therapy or surgical removal of tumor tissue, frequently do not result in a complete cure.

The cancer phenomenon often goes along with overexpression or underexpression of certain genes in degenerated cells, it still being unclear whether these altered expression rates are the cause or the result of the malignant transformation.

Identification of these genes would be an important step for development of new therapies against cancer. Spontaneous formation of cancer is often preceded by a host of mutations.

They can have the most varied effects on the expression pattern in the affected tissue, such as, e.g., underexpression or overexpression, but also expression of shortened genes. Several such changes due to these mutation cascades can ultimately lead to malignant degeneration. The complexity of these relationships makes an experimental approach very difficult.

A database that consists of so-called ESTs is used to look for candidate genes, i.e., genes that compared to the tumor tissue are more strongly expressed in normal tissue. (expressed sequence tags) are sequences of cDNAs, i.e., mRNAs transcribed in reverse, therefore molecules that reflect gene expression. The EST sequences are determined for normal and degenerated tissue. These databases are offered to some extent commercially by various companies. The ESTs of the LifeSeq database, which is used here, are generally between 150 and 350 nucleotides long. They represent a pattern that is unmistakable for a certain gene, although this gene is normally very much longer (> 2000 nucleotides). By comparison of the expression patterns of normal and tumor tissue, ESTs can be identified that are important for tumor formation and proliferation. There is, however, the following problem: Since the EST sequences that are found can belong to different regions of an unknown gene due to different constructions of cDNA libraries, in this case a completely incorrect ratio of the occurrence of these ESTs in the respective tissue would arise. This would only be noticed when the complete gene is known and thus ESTs can be assigned to the same gene.

It has now been found that this possibility of error can be reduced if all ESTs from the respective tissue type are assembled beforehand, before the expression patterns are compared to one another. Overlapping ESTs of the same gene were thus combined into longer sequences (see Figure 1, Figure 2a and Figure 3). This lengthening and thus coverage of an essentially larger gene

region in each of the respective bases are intended to largely avoid the above-described error. Since there were no existing software products for this purpose, programs for assembling genomic sections were employed, which were used modified and to which our own programs were added. A flow chart of the assembly procedure is shown in Figures 2b1-2b4.

Nucleic acid sequences Seq. ID No. 1 to Seq. ID No. 31 and Seq. ID 52, which play a role as candidate genes in the hysteromyoma, have now been found.

Nucleic acid sequences Seq. ID Nos. 14-18, 30, 31 and 52 are of special interest.

The invention thus relates to nucleic acid sequences that code a gene product or a portion thereof, comprising

- a) a nucleic acid sequence selected from the group of nucleic acid sequences Seq. ID Nos. 14-18, 30, 31 and 52.
- b) an allelic variation of the nucleic acid sequences named under a)

or

c) a nucleic acid sequence that is complementary to the nucleic acid sequences named under a) or b).

In addition, the invention relates to a nucleic acid sequence according to one of the sequences Seq. ID Nos. 14-18, 30, 31 or a complementary or allelic variant thereof and the nucleic acid sequences thereof, which have 90% to 95% homology to a human nucleic acid sequence.

The invention also relates to nucleic acid sequences Seq. ID No. 1 to Seq. ID No. 31 and Seq. ID 52, which are expressed elevated in the hysteromyoma.

The invention further relates to nucleic acid sequences comprising a portion of the above-mentioned nucleic acid sequences in such a sufficient amount that they hybridize with sequences Seq. ID Nos. 14-18, 30, 31 and 52.

The nucleic acid sequences according to the invention generally have a length of at least 50 to 4500 bp, preferably a length of at least 150 to 4000 bp, especially a length of 450 to 3500 bp.

With the partial sequences Seq. ID Nos. 14-18, 30, 31 and 52 according to the invention, expression cassettes can also be built using current process practice, whereby on the cassette at least one of the nucleic acid sequences according to the invention is combined with at least one control or regulatory sequence generally known to one skilled in the art, such as, e.g., a suitable promoter. The sequences according to the invention can be inserted in a sense or antisense orientation.

A large number of expression cassettes or vectors and promoters which can be used are known in the literature.

Expression cassettes or vectors are defined as:

1. bacterial, such as, e.g., phagescript, pBs, φX174, pBluescript
SK, pBs KS, pNH8a, pNH16a, pNH18a, pNH46a (Stratagene), pTrc99A,
pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia), 2. eukaryotic,
such as, e.g., pWLneo, pSV2cat, pOG44, pXT1, pSG (Stratagene),
pSVK3, pBPV, pMSG, pSVL (Pharmacia).

Control or regulatory sequences are defined as suitable promoters. Here, two preferred vectors are the pKK232-8 and the PCM7 vector. In particular, the following promoters are intended: lacI, lacZ, T3, T7, gpt, lambda P_R , trc, CMV, HSV thymidine-kinase, SV40, LTRs from retrovirus and mouse metallothionein-I.

The DNA sequences located on the expression cassette can code a fusion protein which comprises a known protein and a bioactive polypeptide fragment.

The expression cassettes are likewise the subject matter of this invention.

The nucleic acid fragments according to the invention can be used to produce full-length genes. The genes that can be obtained are likewise the subject matter of this invention.

The invention also relates to the use of the nucleic acid sequences according to the invention and the gene fragments that can be obtained from use.

The nucleic acid sequences according to the invention can be moved with suitable vectors into host cells, in which as the heterologous part, the genetic information which is contained on the nucleic acid fragments and which is expressed is located.

The host cells containing the nucleic acid fragments are likewise the subject matter of this invention.

Suitable host cells are, e.g., prokaryotic cell systems such as $\underline{\text{E. coli}}$ or eukaryotic cell systems such as animal or human cells or yeasts.

The nucleic acid sequences according to the invention can be used in the sense or antisense form.

Production of polypeptides or their fragments is done by cultivation of the host cells according to current cultivation methods and subsequent isolation and purification of the peptides or fragments, likewise using current methods. The invention further relates to nucleic acid sequences, which code at least a partial sequence of a bioactive polypeptide.

This invention further relates to polypeptide partial sequences, so-called ORF (open-reading-frame)-peptides, according to the sequence protocols ORF ID Nos. 32-51 and ORF ID Nos. 53-55.

The invention further relates to the polypeptide sequences that have at least 80% homology, especially 90% homology to the polypeptide partial sequences of ORF ID Nos. 32-51 and ORF ID Nos. 53-55 according to the invention.

The invention also relates to antibodies that are directed against a polypeptide or a fragment thereof and that are coded by the nucleic acids of sequences Seq. ID Nos. 1-31 and Seq. ID No. 52 according to the invention.

Antibodies are defined especially as monoclonal antibodies.

The antibodies according to the invention can be identified by, i.a., a phage display process. These antibodies are also the subject matter of the invention.

The polypeptide partial sequences according to the invention can be used in a phage display process. The polypeptides that are identified with this process and that bind to the polypeptide

partial sequences according to the invention are also the subject matter of the invention.

The nucleic acid sequences according to the invention can also be used in a phage display process.

The polypeptides of sequences Seq. ID Nos. 32-51 and Seq. ID Nos. 53-55 according to the invention can also be used as tools for finding active ingredients against hysteromyoma, which is likewise the subject matter of this invention.

Likewise the subject matter of this invention is the use of nucleic acid sequences according to sequences Seq. ID No. 1 to Seq. ID No. 31 and Seq. ID 52 for expression of polypeptides, which can be used as tools for finding active ingredients against hysteromyoma.

The invention also relates to the use of the found polypeptide partial sequences Seq. ID No. 32 to Seq. ID No. 51 and Seq. ID Nos. 53-55 as pharmaceutical agents in the gene therapy for treatment of hysteromyoma or for the production of a pharmaceutical agent for treatment of hysteromyoma.

The invention also relates to pharmaceutical agents that contain at least one polypeptide partial sequence Seq. ID Nos. 32 to 51 and Seq. ID Nos. 53-55.

The nucleic acid sequences found according to the invention can also be genomic or mRNA sequences.

The invention also relates to genomic genes, their exon and intron structures and their splice variants that can be obtained from cDNAs of sequences Seq. ID Nos. 1-31, and 52, and their use

together with suitable regulatory elements, such as suitable promoters and/or enhancers.

With the nucleic acids according to the invention (cDNA sequences) Seq. ID Nos. 1-31 and 52, genomic BAC, PAC and cosmid libraries are screened, and specifically human clones are isolated via complementary base pairing (hybridization). The BAC, PAC and cosmid clones isolated in this way are hybridized using fluorescence-in-situ hybridization on metaphase chromosomes, and the corresponding chromosome sections on which the corresponding genomic genes lie are identified. BAC, PAC and cosmid clones are sequenced in order to clarify the corresponding genomic genes in their complete structure (promoters, enhancers, silencers, exons and introns). BAC, PAC and cosmid clones can be used as independent molecules for gene transfer (see Figure 5).

The invention also relates to BAC, PAC and cosmid clones containing functional genes and their chromosomal localization according to sequences Seq. ID Nos. 1-31 and 52, for use as vehicles for gene transfer.

Meanings of T chnical Terms and Abbreviations

Nucleic acids = Nucleic acids in this invention are defined as:

mRNA, partial cDNA, full-length cDNA and genomic

genes (chromosomes)

ORF = Open Reading Frame, a defined sequence of amino acids which can be derived from the cDNA sequence

Contig = A set of DNA sequences that can be combined as a result of very great similarities into one sequence (consensus)

Singleton = A contig that contains only one sequence

Module = Domain of a protein with a defined sequence, which represents one structural unit and which occurs in various proteins

N = selectively the nucleotide A, T, G or C

X = selectively one of the 20 naturally occurring
amino acids

Explanation of the Alignment Parameters

minimal initial match = minimal initial identity area
maximum pads per read = maximum number of insertions
maximum percent mismatch = maximum deviation in %

Explanation of Figures

Figure 1 shows the systematic gene search in the Incyte LifeSeq database

Figure 2a shows the principle of EST assembling
Figures 2b1-2b4 show the entire principle of EST assembling

Figure 3 shows the in-silico subtraction of gene expression in various tissues

Figure 4a shows the determination of tissue-specific expression via electronic Northern

Figure 4b shows the electronic Northern

Figure 5 shows the isolation of genomic BAC and PAC clones.

The following examples explain the production of the nucleic acid sequences according to the invention without limiting the invention to these examples and nucleic acid sequences.

Example 1

Search for Tumor-related Candidate Genes

First, all ESTs of the corresponding tissue from the LifeSeq database (from October 1997) were extracted. They were then assembled by means of the GAP4 program of the Staden package with the parameters 0% mismatch, 8 pads per read and a minimal match The sequences (fails) not recorded in the GAP4 database were assembled first at 1% mismatch and then again at 2% mismatch with the database. Consensus sequences were computed from the contigs of the database that consisted of more than one sequence. The singletons of the database, which consisted of only one sequence, were re-assembled at 2% mismatch with the sequences not recorded in the GAP4 database. In turn, the consensus sequences were determined for the contigs. All other ESTs were reassembled at 4% mismatch. The consensus sequences were extracted once again and finally assembled with the previous consensus sequences and the singletons and the sequences not recorded in the database at 4% mismatch. The consensus sequences were formed and used with the singletons and fails as the initial basis for tissue comparisons. This procedure ensured that among the parameters used, all sequences represented gene regions independent of one another.

Figures 2b1-2b4 illustrate the lengthening of the hysteromyomic tissue ESTs.

The sequences of the respective tissue assembled in this way were then compared to one another by means of the same program (Figure 3). To do this, first all sequences of the first tissue were input into the database. (It was therefore important that they were independent of one another.)

Then, all sequences of the second tissue were compared to all those of the first. The result was sequences that were specific to the first or the second tissue as well as those which occurred in both. In the latter, the ratio of the frequency of occurrence in the respective tissue was evaluated. All programs pertaining to the evaluation of the assembled sequences were themselves developed.

All sequences that occurred more than four times in respectively one of the compared tissues and all that occurred at least five times as often in one of the two tissues were further studied. These sequences were subjected to an electronic Northern (see Example 2.1), by which the distribution in all tumor and normal tissues was studied (see Figure 4a and Figure 4b). The relevant candidates were then lengthened using all Incyte ESTs and all ESTs of public databases (see Example 3). Then, the sequences and their translation into possible proteins were compared to all nucleotide and protein databases and were studied for possible regions that code for proteins.

Example 2

Algorithm for Identification and Lengthening of Partial cDNA Sequences with Altered Expression Pattern

An algorithm for finding overexpressed or underexpressed genes will be explained below. The individual steps are also summarized in a flow chart for the sake of clarity (see Figure 4b).

2.1. Electronic Northern Blot

By means of a standard program for homology search, e.g.,
BLAST (Altschul, S. F.; Gish, W.; Miller, W.; Myers, E. W. and
Lipman, D. J. (1990) J. Mol. Biol. 215, 403-410), BLAST2

(Altschul, S. F.; Madden, T. L.; Schäffer, A. A.; Zhang, J.;
Zhang, Z.; Miller, W., and Lipman, D. J. (1997) Nucleic Acids
Research 25 3389-3402) or FASTA (Pearson, W. R. and Lipman, D. J.

(1988) Proc. Natl. Acad. Sci. USA 85 2444-2448), the homologous
sequences in various EST libraries (private or public) arranged
by tissues are determined for a partial DNA sequence S, e.g., an
individual EST or a contig of ESTs. The (relative or absolute)
tissue-specific occurrence frequencies of this partial sequence S
which were determined in this way are called electronic Northern
Blots.

2.1.1

Analogously to the procedure described under 2.1, the sequence Seq. ID No. 30 was found, which occurs more frequently in the hysteromyomic tissue than in normal tissue.

The result is as follows:

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Electronic Northern for SEQ. ID NO.: 30

	NO	ORMAL	TU	JMOR	Rati	.os
	ક	frequency	કૃ	frequency	N/T	T/N
		0.0070		0.0000	undef	0.0000
Bladder		0.0078		0.0000	undef	0.0000
Breast		0.0013 0.0092		0.0000	undef	
Small intestine		0.0092		0.0000	undef	
Ovary		0.0000		0.0000	undef	
Endocrine tissue		0.0019		0.0000	undef	
Gastrointestinal		0.0000		0.0000	undef	
Brain		0.0000		0.0000	undef	
Hematopoietic		0.0000		0.0000	undef	
Skin		0.0000		0.0000	undef	
Hepatic		0.0011		0.0000		0.0000
Heart		0.0000		0.0000	undef	
Testicles		0.0010		0.0000	undef	0.0000
Lung		0.0000		0.0000	undef	undef
Stomach-esophagus		0.0000		0.0000	undef	undef '
Muscle-skeleton		0.0000		0.0000	undef	undef
Kidney		0.0000	·	0.0000	undef	undef
Pancreas		0.0000		0.0000	undef	undef
Penis		0.0109		0.0021		0.1954
Prostate		0.0000		0.0000	undef	
Uterus-endometrium		0.0000		0.0340	0.0000	
Uterus-myometrium		0.0102		0.0000	undef	0.0000
Uterus-general		0.0000	•			
Breast hyperplasia		0.0000				
Prostate hyperplasia		0.0089			•	
Seminal vesicle		0.0000				
Sensory organs		0.0000				
White blood cells		0.0000				
Cervix				,		
CCIVIA						

. · · · · · · · · · · · · · · · · · · ·	FETUS % frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000

2.1.2.

Analogously to the procedure described under 2.1, the sequence Seq. ID No. 31 was found, which occurs more frequently in the hysteromyomic tissue than in normal tissue.

The result is as follows:

Electronic Northern for SEQ. ID NO.: 31

	NORMAL % freque	TUMOR ency % freque	Ratios ency N/T	T/N
Bladder Breast Small intestine Ovary Endocrine tissue Gastrointestinal Brain Hematopoietic Skin Hepatic Heart Testicles Lung Stomach-esophagus Muscle-skeleton Kidney Pancreas Penis Prostate Uterus-endometrium Uterus-myometrium Uterus-general Breast hyperplasia Prostate hyperplasia Seminal vesicle Sensory organs	0.0000 0.0038 0.0031 0.0030 0.0034 0.0057 0.0111 0.0040 0.0000 0.0042 0.0058 0.0042 0.0000 0.0034 0.0163 0.0017 0.0090 0.0022 0.0135 0.0000 0.0022 0.0000 0.0022 0.0000 0.0022	0.0051 0.0019 0.0000 0.0104 0.0100 0.0046 0.0051 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0060 0.0068 0.0068 0.0000 0.0000 0.0000 0.0000 0.0000	0.0000 undef 2.0416 0.4898 undef 0.0000 0.2878 3.4745 0.3396 2.9444 1.2425 0.8048 2.1599 0.4630 undef 0.0000 undef undef undef 0.0000 undef 0.0000 undef 0.0000 undef 0.0000 undef 0.0000 undef 0.0000 undef 0.0000 0.4064 2.4605 undef undef 0.5711 1.7510 2.3791 0.4203 undef 0.0000 undef 0.0000 undef 0.0000 undef 0.0000	T/N
White blood cells Cervix	0.0213			

	FETUS % frequency	STANDARDIZED/SUBTRACTED LIBRARIES % frequency		
Development Gastrointestinal Brain Hematopoietic Skin Hepatic	0.0000 0.0028 0.0125 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal	0.0000 0.0000 0.0000 0.0000 0.0000	
Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0036 0.0036 0.0254 0.0062 0.0303 0.0000	Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0000 0.0000 -0.0082 0.0000 0.0000	

In an analogous procedure, the following Northerns were also found:

Electronic Northern			_
	NORMAL	TUMOR	Ratios
	% frequency	<pre>% frequency</pre>	N/T T/N
Bladder	0.0468	0.0562	0.8321 1.2018
Breast	0.0576	0.0752	0.7656 1.3062
Small intestine	0.0766	0.0662	1.1586 0.8631
Ovary	0.0509	0.0650	0.7829 1.2774
Endocrine tissue	0.0596	0.0702	0.8491 1.1778
Gastrointestinal	0.0690	0.1203	0.5735 1.7438
Brain	0.0850	0.0873	0.9741 1.0266 1.9056 0.5248
Hematopoietic	0.0722	0.0379	undef 0.0000
Skin	0.0918	0.0000	0.1298 7.7066
Hepatic	0.0143	0.1100	4.0862 0.2447
Heart	0.1123	0.0275 0.0819	0.6325 1.5809
Testicles	0.0518 0.1080	0.0879	1.2287 0.8138
Lung	0.0676	0.0307	2.2059 0.4533
Stomach-esophagus	0.0737	0.0780	0.9445 1.0587
Muscle-skeleton	0.0407	0.0890	0.4575 2.1857
Kidney	0.0463	0.0552	0.8376 1.1939
Pancreas	0.0449	0.0000	undef 0.0000
Penis	0.0436	0.0554	0.7874 1.2700
Prostate	0.0338	0.0000	undef 0.0000
Uterus-endometrium	0.0534	0.1426	0.3741 2.6732
	0.0255	0.0000	undef 0.0000
Uterus-myometrium	0.0863		
Uterus-general	0.0535		
Breast hyperplasia	0.0890		
Prostate hyperplasia			
Seminal vesicle	0.0772		
Sensory organs	0.1171		
White blood cells			
Cervix			

*	FETUS % frequency	STANDARDIZED/SUBTRA LIBRARIES % frequency	CTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0139 0.0583 0.0500 0.0590 0.0000 0.0000 0.0462 0.0578 0.0254 0.0679 0.0545 0.2493 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus_n	0.0204 0.1595 0.0051 0.0256 0.0488 0.0000 0.0227 0.0154 0.0082 0.0191 0.0068 0.0000 0.0000

Electronic Northern	for SEQ. ID NO NORMAL	O.: 2 TUMOR	Ratios
	% frequency	% frequency	N/T T/N
Bladder			
Breast	0.0039	0.0102	0.3814 2.6222
Small intestine	0.0128	0.0038	3.4026 0.2939
	0.0031	0.0165	0.1854 5.3946
Ovary	0.0000	0.0000	undef undef
Endocrine tissue	0.0017	0.0075	0.2264 4.4166
Gastrointestinal	0.0019	0.0093	0.2071 4.8289
Brain	0.0022	0.0103	0.2160 4.6299
Hematopoietic	0.0040	0.0000	undef 0.0000
Skin	0.0000	0.0000	undef undef
Hepatic	0.0000	0.0065	0.0000 undef
Heart	0.0011	0.0137	0.0771 12.9706
Testicles	0.0058	0.0000	undef 0.0000
Lung	0.0021	0.0082	0.2540 3.9367
Stomach-esophagus	0.0000	0.0000	undef undef
Muscle-skeleton	0.0000	0.0120	0.0000 undef
Kidney	0.0054	0.0000	undef 0.0000
Pancreas	0.0000	0.0055	0.0000 undef
Penis	0.0030	0.0000	undef 0.0000
Prostate	0.0000	0.0021	0.0000 undef
Uterus-endometrium	0.0000	0.0528	0.0000 undef
Uterus-myometrium	0.0000	0.0408	0.0000 undef 0.0534 18.7357
Uterus-general	0.0051 0.0064	0.0954	0.0334 18.7337
Breast hyperplasia	0.0000		
Prostate hyperplasia		•	
Seminal vesicle	0.0000		
Sensory organs	0.0009		
White blood cells	0.0000		
Cervix	3.3300		
CETATY			

	FETUS % frequency	STANDARDIZED/SUBTRACTED LIBRARIES & frequency)
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate	0.0063 0.0157 0.0000 0.0000 0.0249 0.0325 0.0761 0.0618 0.0727 0.0249	Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate	.0000 .0000 .0000 .0384 .0122 .0000 .0259 .0000 .0328 .0171 .0137
Sensory organs	0.0000	Sensory Organs 0 Uterus_n	.0083

Electronic Northern	for SEQ. ID No NORMAL % frequency	TUMOR	Ratios cy N/T	T/N
Bladder Breast Small intestine Ovary Endocrine tissue Gastrointestinal Brain Hematopoietic Skin Hepatic Heart Testicles Lung Stomach-esophagus Muscle-skeleton Kidney Pancreas Penis Prostate Uterus-endometrium Uterus-myometrium Uterus-general Breast hyperplasia Prostate hyperplasia Prostate hyperplasia Seminal vesicle Sensory organs White blood cells Cervix	0.0000 0.0000	0.0000 0.0000 0.0000 0.1103 0.0000 0.0010 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	undef	0.0731

	FETUS % frequency	LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0000 0.0028 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000
		Uterus_n	

Electronic Northern fo	or SEQ. ID NO	0.: 4	
	NORMAL	TUMOR	Ratios
	<pre>% frequency</pre>	% frequenc	y N/T T/N
Bladder	0.0078	0.0077	1.0170 0.9833
Breast	0.0026	0.0038	0.6805 1.4694
Small intestine	0.0031	0.0165	0.1854 5.3946
Ovary	0.0060	0.0026	2.3025 0.4343
Endocrine tissue	0.0000	0.0000	undef undef
Gastrointestinal	0.0000	0.0046	0.0000 undef
Brain	0.0007	0.0031	0.2400 4.1669
Hematopoietic	0.0000	0.0000	undef undef
Skin	0.0000	0.1695	0.0000 undef
Hepatic	0.0000 0.0095	0.0065	0.0000 undef
Heart	0.0000	0.0412	0.2313 4.3235
Testicles	0.0021	0.0000 0.0041	undef undef
Lung	0.0000	0.0077	0.5080 1.9684
Stomach-esophagus	0.0017	0.0120	0.0000 undef
Muscle-skeleton	0.0027	0.0000	0.1428 7.0040 undef 0.0000
Kidney	0.0033	0.0055	0.5983 1.6714
Pancreas	0.0030	0.0000	undef 0.0000
Penis	0.0000	0.0021	0.0000 undef
Prostate	0.0068	0.0000	undef 0.0000
Uterus-endometrium	0.0152	0.0883	0.1727 5.7919
Uterus-myometrium	0.0051	0.0000	undef 0.0000
Uterus-general	0.0032		
Breast hyperplasia	0.0030		
Prostate hyperplasia	0.0000		
Seminal vesicle	0.0118 0.0000		
Sensory organs	0.0000		
White blood cells	0.000		
Cervix			

	FETUS % frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0278 0.0139 0.0000 0.0000 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0068 0.0000 0.0000 0.0000 0.0093 0.0122 0.0000
Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0253 0.0000 0.0185 0.0061 0.0000 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs	0.0130 0.0000 0.0000 0.0000 0.0000 0.0000

Electronic Northern		0.: 5	
	NORMAL	TUMOR	Ratios
	% frequency	% frequenc	y N/T T/N
		-	
Bladder			
Breast	0.0390	0.0332	1.1734 0.8522
Small intestine	0.0345	0.0414	0.8352 1.1973
Ovary	0.0399	0.0662	0.6024 1.6599
Endocrine tissue	0.0359	0.0546	0.6579 1.5201
Gastrointestinal	0.0528	0.0351	1.5040 0.6649
Brain	0.0172	0.0185	0.9319 1.0731
Hematopoietic.	0.0214	0.0359	0.5965 1.6763
Skin	0.0294	0.0379	0.7763 1.2881
Hepatic	0.0257	0.1695	0.1516 6.5954
Heart	0.0476	0.0323	1.4706 0.6800
Testicles	0.0276	0.0000	undef 0.0000
	0.0633	0.0234	2.7059 0.3696
Lung	0.0312	0.0266	1.1724 0.8530
Stomach-esophagus	0.0387 0.0308	0.0153 0.0360	2.5211 0.3967
Muscle-skeleton	0.0306	0.0548	0.8567 1.1673 0.5948 1.6813
Kidney	0.0132	0.0221	0.5983 1.6714
Pancreas	0.0479	0.0000	undef 0.0000
Penis	0.0153	0.0319	0.4777 2.0934
Prostate	0.0473	0.0528	0.8962 1.1158
Uterus-endometrium	0.0305	0.1019	0.2993 3.3415
Uterus-myometrium	0.0560	0.1908	0.2936 3.4065
Uterus-general	0.0448		
Breast hyperplasia	0.0446		
Prostate hyperplasia	0.1513		
Seminal vesicle	0.0235	,	
Sensory organs	0.0061		
White blood cells	0.0426		•
Cervix			

	FETUS % frequency	LIBRARIES & frequency	
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0974 0.0222 0.0000 0.0197 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0136 0.0000 0.0051 0.0000 0.0099 0.0122 0.0000
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0325 0.0000 0.0185 0.0909 0.1496 0.0126	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0231 0.0328 0.0131 0.0068 0.0077 0.0208

STANDARDIZED/SUBTRACTED

Electronic Northern	for SEQ. ID NO.: 6 NORMAL TUMON % frequency % fre	
Bladder	0.0429 0.	1278 0.3356 2.9798
Breast		1203 0.8613 1.1610
Small intestine		0.2163 1.1610
Ovary		0702 0.8528 1.1726
Endocrine tissue		0.426 1.8380 0.5441
Gastrointestinal		0786 0.3167 3.1574
Brain		1284 0.3341 2.9935
Hematopoietic		1136 0.2000 5.0008
Skin		1695 0.4982 2.0073
Hepatic		0712 0.7353 1.3600
Heart		1649 0.5590 1.7890
Testicles		0585 0.7872 1.2704
		0797 0.5601 1.7853
Lung Stomach-esophagus		0537 0.0000 undef
Muscle-skeleton		1380 1.0801 0.9258
	0.0489 0.	0479 1.0196 0.9808
Kidney	0.0149 0.	1049 0.1417 7.0571
Pancreas	0.0509 0.	1333 0.3819 2.6187
Penis		0.6142 1.6282
Prostate		1583 0.4268 2.3432
Uterus-endometrium		3260 0.1169 8.5541
Uterus-myometrium		0000 undef 0.0000
Uterus-general	0.0863	
Breast hyperplasia	0.0297	
Prostate hyperplasia	0.0089	
Seminal vesicle	0.0353	
Sensory organs	0.0078	
White blood cells	0.0213	•
Cervix		
	CTAI	IDADDIZED/SIBTDACTED

	FETUS % frequency	LIBRARIES % frequency	CTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels		Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0476 0.1595 0.0203 0.0000 0.0379 0.0122 0.0000
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0889 0.0759 0.2535 0.1112 0.1091 0.1247 0.1004	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs	0.0486 0.0077 0.0164 0.0251 0.0068 0.0155 0.0167

Electronic Northern	for SEQ. ID NO NORMAL	O.: 7 TUMOR	Ratios
		% frequency	
Bladder	0.0117	0.0051	2.2882 0.4370
Breast	0.0117	0.0031	1.7013 0.5878
Small intestine	0.0153	0.0331	0.4634-2.1579
Ovary	0.0150	0.0104	1.4391 0.6949
Endocrine tissue	0.0085	0.0050	1.6981 0.5889
Gastrointestinal	0.0096	0.0278	0.3451 2.8974
Brain	0.0059	0.0113	0.5236 1.9098
Hematopoietic	0.0254	0.0000	undef 0.0000
Skin	. 0.0257	0.0000	undef 0.0000
Hepatic	0.0143	0.0194	0.7353 1.3600
Heart	0.0106	0.0000	undef 0.0000
Testicles	0.0058	0.0234	0.2460 4.0652
Lung	0.0104	0.0245	0.4234 2.3620
Stomach-esophagus	0.0000	0.0000	undéf undef
Muscle-skeleton	0.0051	0.0360	0.1428 7.0040
	0.0244	0.0068	3.5687 0.2802
Kidney	0.0083	0.0000	undef 0.0000
Pancreas	0.0180	0.0000	undef 0.0000
Penis	. 0.0218 . 0.0135	0.0064 0.0000	3.4121 0.2931 undef 0.0000
Prostate	0.0000	0.0408	0.0000 undef
Uterus-endometrium	0.0000	0.0000	undef undef
Uterus-myometrium	0.0192		ander under
Uterus-general	0.0030	•	
Breast hyperplasia	0 0000	ii. •	
Prostate hyperplasia	0.0235		
Seminal vesicle	0.0208		
Sensory organs	0.0213		
White blood cells			
Cervix		•	÷
			•
			ED/SUBTRACTED
	FETUS % frequency	LIBRARIES % frequence	
Development		Breast	0.000

	% frequency	% frequency	
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0278 0.0083 0.0125 0.0000 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0000 0.0000 0.0101 0.0245 0.0093 0.0122 0.0342
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0036 0.0000 0.0309 0.0061 0.0249 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0097 0.0154 0.0082 0.0080 0.0000 0.0000

Electronic Northern fo	or SEQ. ID NO NORMAL	D.: 8 TUMOR	Ratios
	<pre>% frequency</pre>	% frequency	N/T T/N
Bladder	0.0312	0.0460	
Breast	0.0192	0.0460 0.0282	0.6780 1.4750
Small intestine	0.0368	0.0000	0.6805 1.4694
	0.0210	0.0364	undef 0.0000 0.5756 1.7372
Ovary	0.0290	0.0326	0.8882 1.1258
Endocrine tissue	0.0460	0.0231	1.9880 0.5030
Gastrointestinal	0.0532	0.0585	0.9094 1.0996
Brain	0.0361	0.0379	0.9528 1.0496
Hematopoietic	0.0367	0.0000	undef 0.0000
Skin	0.0048	0.0647	0.0735 13.5999
Hepatic	0.0699	0.0412	1.6961 0.5896
Heart	0.0288	0.4210	0.0683 14.6349
Testicles	0.0343	0.0368	0.9314 1.0737
Lung	0.0773	0.0230	3.3614 0.2975
Stomach-esophagus	0.0497 0.0353	0.0660	0.7528 1.3283
Muscle-skeleton	0.0165	0.1575	0.2241 4.4619
Kidney	0.0299	0.0939 0.0267	0.1760 5.6828
Pancreas	0.0196	0.0298	1.1232 0.8903
Penis	0.0270	0.1583	0.6580 1.5197 0.1707 5.8579
Prostate	0.0229	0.0679	0.3367 2.9702
Uterus-endometrium	0.0051	0.0954	0.0534 18.7357
Uterus-myometrium	0.0192		0.0334 20.7337
Uterus-general	0.0505		
Breast hyperplasia	0.0890		
Prostate hyperplasia	0.0353	•	=
Seminal vesicle	0.0399		
Sensory organs	0.0319	•	*
White blood cells		•	•
Cervix			

	FETUS % frequency	STANDARDIZED/SUBTRACTED LIBRARIES % frequency	
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0417 0.0333 0.0313 0.0197 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0000 0.0000 0.0152 0.0000 0.0082 0.0244 0.0057 0.0032
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0217 0.0507 0.0309 0.0727 0.0997 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0077 0.0082 0.0141 0.0000 0.0310 0.0125

Electronic Northern f	or SEQ. ID NO.	: 9	
		'UMOR	Ratios
	% frequency %	frequency	N/T T/N
			•
Bladder	0.0468	0.0204	2 2002 0 4170
Breast	0.0205	0.0204	2.2882 0.4370
Small intestine	0.0203	0.0431	0.4537 2.2042
Ovary	0.0539	0.0351	0.9268 1.0789 1.1513 0.8686
Endocrine tissue	0.0562	0.0468	1.1513 0.8686
Gastrointestinal	0.0362	0.0327	0.7692 1.3001
Brain	0.0333	0.0318	1.0451 0.9568
	0.0388	0.0318	1.0234 0.9772
Hematopoietic	0.0257	0.1695	0.1516 6.5954
Skin	0.0095	0.0259	0.3676 2.7200
Hepatic	0.0540	0.0000	undef 0.0000
Heart	0.0173	0.0702	0.2460 4.0652
Testicles	0.0291	0.0491	0.5927 1.6872
Lung	0.0000	0.0230	0.0000 undef
Stomach-esophagus	0.0206	0.1140	0.1803 5.5448
Muscle-skeleton	0.0733	0.0274	2.6765 0.3736
Kidney	0.0264	0.0331	0.7977 1.2536
Pancreas	0.0269	0.0533	0.5054 1.9786
Penis	0.0501	0.0341	1.4715 0.6796
Prostate	0.0541	0.2639	0.2049 4.8816
Uterus-endometrium	0.0152	0.0611	0.2494 4.0097
Uterus-myometrium	0.0204	0.0000	under 0.0000
Uterus-general	0.0512		
Breast hyperplasia	0.0386		
Prostate hyperplasia	0.0623		
Seminal vesicle	0.0470		
Sensory organs	0.0286		
White blood cells	0.0426	*	•
Cervix			
CCTATY			

	FETUS % frequency	STANDARDIZED/SUBTRACTEI LIBRARIES % frequency	
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0557 0.0666 0.0626 0.0786 0.0000 0.0260	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0068 0.1595 0.0051 0.0245 0.0256 0.0732 0.0057
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0818 0.0867 0.0761 0.0432 0.0606 0.0249 0.0628	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0309 0.0737 0.0231 0.0137 0.0310 0.0333

Electronic Northern	for SEQ. ID NORMAL	O.: 10 TUMOR	Ratios
	% frequency	% frequenc	y n/T T/n
Bladder	A	-	
Breast	0.1131	0.1431	0.7900 1.2659
	0.1356	0.1692	0.8015 1.2476
Small intestine	0.1962 0.1827	0.0165	11.8636 0.0843
Ovary	0.1090	0.2446 0.1329	0.7471 1.3385
Endocrine tissue	0.1090	0.1329	0.8202 1.2192
Gastrointestinal	0.1035	0.2390	0.7248 1.3797
Brain	0.2700	0.1323	0.7814 1.2798 1.4257 0.7014
Hematopoietic	0.1358	0.0847	1.6028 0.6239
Skin	0.0428	0.1812	0.2363 4.2311
Hepatic	0.2586	0.0137	18.8118 0.0532
Heart	0.0633	0.1403	0.4510 2.2174
Testicles	0.3231	0.2229	1.4495 0.6899
Lung	0.1643	0.2147	0.7653 1.3066
Stomach-esophagus	0.1970	0.0960	2.0524 0.4872
Muscle-skeleton	0.1222	0.2328	0.5248 1.9055
Kidney	0.0876	0.1767	0.4955 2.0183
Pancreas	0.1407	0.1600	0.8798 1.1366
Penis	0.1003	0.0766	1.3080 0.7645
Prostate	0.1824	0.0528	3.4569 0.2893
Uterus-endometrium	0.1677	0.3804	0.4409 2.2681
Uterus-myometrium	0.2292	0.0000	undef 0.0000
Uterus-general	0.0735		
Breast hyperplasia	0.0684		
Breast hyperprasia	0.1335		
Prostate hyperplasia Seminal vesicle			•
	0.2749		•
Sensory organs	0.0958		•
White blood cells			
Cervix			

	FETUS % frequency	STANDARDIZED/SUBTRA LIBRARIES % frequency	CTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0139 0.0639 0.1063 0.1258 0.2513 0.0260 0.0712	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle	0.0136 0.3190 0.0709 0.0000 0.0326 0.1464 0.0000 0.1328
Suprarenal gland Kidney Placenta Prostate Sensory organs	0.1517 0.0000 0.0988 0.0909 0.1745 0.0377	Testicles Lung Nerves Prostate Sensory Organs	0.2211 0.0311 0.0410 0.0000 0.0333

Electronic Northern	for SEQ. ID No	0.: 11	•
	NORMAL	TUMOR	Ratios
	% frequency	% frequence	cy N/T T/N
		-	- ,
Bladder			•
Breast	0.0156	0.0051	3.0509 0.3278
Small intestine	0.0064	0.0113	0.5671 1.7633
Ovary	0.0031	0.0000	undef 0.0000
Endocrine tissue	0.0240	0.0234	1.0233 0.9772
Gastrointestinal	0.0068	0.0100	0.6792 1.4722
Brain	0.0096	0.0046	2.0708 0.4829
	0.0022	0.0257	0.0864 11.5747
Hematopoietic	0.0147	0.0000	undef 0.0000
Skin	0.0110	0.0000	undef 0.0000
Hepatic	0.0238	0.0065	3.6765 0.2720
Heart	0.0276	0.0275	1.0023 0.9977
Testicles	0.0000	0.0117	0.0000 undef
Lung	0.0083	0.0245	0.3387 2.9526
Stomach-esophagus	0.0000	0.0077	0.0000 undef
Muscle-skeleton	0.0086	0.0060	1.4278 0.7004
Kidney	0.0081	0.0616	0.1322 7.5658
Pancreas	0.0165	0.0000	undef 0.0000
Penis	0.0120	0.0267	0.4493 2.2259
Prostate	0.0065	0.0085	0.7677 1.3026
Uterus-endometrium	0.0270	0.0000	undef 0.0000
	0.0076	0.0951	0.0802 12.4748
Uterus-myometrium	0.0968	0.0000	undef 0.0000
Uterus-general	0.0096		
Breast hyperplasia	0.0089		
Prostate hyperplasia			-
Seminal vesicle	0.0706	•	
Sensory organs	0.0000		
White blood cells	0.0106		
Cervix			

	FETUS % frequency	LIBRARIES % frequency	CACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung	0.0139 0.0278 0.0000 0.0708 0.0000 0.0000 0.0320	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle	0.0000 0.0000 0.0000 0.0000 0.0233 0.0244 0.0000 0.0648
Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0000 0.0124 0.2121 0.0000 0.0000	Testicles Lung Nerves Prostate Sensory Organs Uterus_n	0.0000 0.0164 0.0010 0.0068 0.0000 0.0083

Electronic Northern			
		UMOR	Ratios
	<pre>% frequency %</pre>	frequency	N/T T/N
		,	
Bladder	2 2353		0 1135 0 0136
Breast	0.0351 0.0269	0.3093	0.1135 8.8135
Small intestine		0.0470	0.5716 1.7493
Ovary	0.0092	0.0662	0.1390 7.1929
Endocrine tissue	0.0569 0.0528	0.0182	3.1248 0.3200
	0.0328	0.0502	1.0528 0.9498
Gastrointestinal	0.0022	0.0093	0.2071 4.8289
Brain	0.0022	0.0873 0.0000	0.0254 39.3541 undef 0.0000
Hematopoietic	0.0220	0.0000	0.0433 23.0839
Skin	0.0285	0.0582	0.4902 2.0400
Hepatic	0.0191	0.0000	undef 0.0000
Heart	0.0191	0.0000	undef 0.0000
Testicles	0.0113	0.0061	1.8628 0.5368
Lung	0.0193	0.0077	2.5211 0.3967
Stomach-esophagus	0.0771	0.0540	1.4278 0.7004
Muscle-skeleton	0.0489	0.0137	3.5687 0.2802
Kidney	0.0264	0.0442	0.5983 1.6714
Pancreas	0.0090	0.1066	0.0842 11.8713
Penis	0.0000	0.0064	0.0000 undef
Prostate	0.0743	0.0000	undef 0.0000
Uterus-endometrium	0.0381	0.1494	0.2551 3.9206
Uterus-myometrium	0.0153	0.0954	0.1601 6.2452
Uterus-general	0.0096		•
Breast hyperplasia	0.0059		
Prostate hyperplasia	0.0000		
Seminal vesicle	0.0118		
Sensory organs	0.0000		
White blood cells	0.0000		
Cervix			
CELAIX			

	FETUS % frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0696 0.4387 0.0000 0.2713 0.0000 1.6121 0.0605 0.1770 1.1663 0.8092 0.7635 0.0499 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus_n	0.0612 0.0000 0.1114 0.0000 0.4665 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0068 0.0000 0.0291

Electronic Northern			D-+1
	NORMAL	TUMOR	Ratios
	<pre>% frequency</pre>	% frequency	N/T T/N
Bladder	·	•	
Breast	0.0078	0.0204	0.3814 2.6222
Small intestine	0.0256	0.0432	0.5918 1.6899
Ovary	0.0000	0.0000	undef undef
Endocrine tissue	0.0060	0.0078	0.7675 1.3029
Gastrointestinal	0.0000	0.0050	0.0000 undef
Brain	0.0000	0.0000	undef undef
Hematopoietic	0.0074 0.0013	0.0216 0.0000	0.3428 2.9168 undef 0.0000
Skin	0.0013	0.0000	under 0.0000 under 0.0000
Hepatic	0.0000	0.0000	under under
Heart	0.0021	0.0000	undef 0.0000
Testicles	0.0115	0.0468	0.2460 4.0652
Lung	0.0031	0.0082	0.3810 2.6245
Stomach-esophagus	0.0580	0.0000	undef 0.0000
Muscle-skeleton	0.0000	0.0300	0.0000 undef
Kidney	0.0027	0.0000	undef 0.0000
Pancreas	0.0033	0.0276	0.1197 8.3571
Penis	0.0210	0.1066	0.1966 5.0877
Prostate	0.0022	0.0043	0.5118 1.9538
	0.0135	0.0000	undef 0.0000
Uterus-endometrium	0.0305	0.0883	0.3453 2.8959
Uterus-myometrium	0.0102	0.0000	undef 0.0000
Uterus-general	0.0160	•	
Breast hyperplasia	0.0059		
Prostate hyperplasia	0.0356	• •	
Seminal vesicle	0.0000		
Sensory organs	0.0035 0.0639		
White blood cells	0.0639		
Cervix			

	FETUS % frequency	STANDARDIZED/SUBT LIBRARIES % frequency	RACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0000 0.0028 0.0000 0.0000 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0408 0.0000 0.1266 0.0000 0.0047 0.0000 0.0057 0.0032
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0145 0.0761 0.0247 0.0061 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0000 0.0000 0.0030 0.0000 0.0000 0.0208

Electronic Northern			
		UMOR	Ratios
	<pre>% frequency %</pre>	frequency	N/T T/N
Bladder	•		
Breast	0.0195	0.0051	3.8136 0.2622
Small intestine	0.0051	0.0132	0.3889 2.5715
Ovary	0.0061	0.0000	undef 0.0000
Endocrine tissue	0.0150	0.0130	1.1513 0.8686
	0.0153	0.0150	1.0189 0.9815
Gastrointestinal	0.0115	0.0000	undef 0.0000
Brain	0.0133	0.0133	0.9969 1.0031
Hematopoietic	0.0174	0.0000	undef 0.0000
Skin	0.0147	0.0000	undef 0.0000
Hepatic	0.0000	0.0194	0.0000 undef
Heart	0.0265	0.0000	undef 0.0000
Testicles	0.0058	0.0000	undef 0.0000
Lung	0.0042	0.0164	0.2540 3.9367
Stomach-esophagus	0.0000	0.0077	0.0000 undef
Muscle-skeleton	0.0069	0.0540	0.1269 7.8795
Kidney	0.0217	0.0274	0.7930 1.2610
Pancreas	0.0165	0.0055	2.9915 0.3343
Penis	0.0090 0.0087	0.0000	undef 0.0000
Prostate	0.0338	0.0170 0.2111	0.5118 1.9538
Uterus-endometrium	0.0076	0.0475	0.1600 6.2484 0.1603 6.2374
Uterus-myometrium	0.0000	0.0000	undef undef
Uterus-general	0.0320	0.000	dider dider
Breast hyperplasia	0.0238		
Prostate hyperplasia			•
Seminal vesicle	0.0000	,	
Sensory organs	0.0069		
White blood cells	0.0213		
Cervix			
OCT 112			

	FETUS % frequency	STANDARDIZED/SUBTE LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0417 0.0139 0.0125 0.0157 0.0000 0.0000 0.0427 0.0253 0.0000 0.0124 0.0121 0.0000 0.0126	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0068 0.0000 0.0051 0.0245 0.0035 0.0244 0.0000 0.0065 0.0000 0.0082 0.0050 0.0137 0.0155 0.0042

Electronic Northern			
	NORMAL	TUMOR	Ratios
	<pre>% frequency</pre>	% frequency	N/T T/N
Bladder	0.0078	0.0102	0.7627 1.3111
Breast	0.0078	0.0150	0.1701 5.8778
Small intestine	0.0028	0.0130	0.3707 2.6973
Ovary	0.0000	0.0052	0.0000 undef
Endocrine tissue	0.0000		0.0000 under
Gastrointestinal	0.0000	0.0023	0.0000 undef
Brain	0.0007	0.0072	0.1029 9.7228
Hematopoietic	0.0000	0.0000	undef undef
Skin	0.0037	0.0000	undef 0.0000
Hepatic	0.0000	0.0000	undef undef
Heart	0.0032	0.0000	undef 0.0000
Testicles	0.0058	0.0000	undef 0.0000
Lung	0.0021	0.0041	0.5080 1.9684
Stomach-esophagus	0.0000	0.0307	0.0000 undef
Muscle-skeleton	0.0086	0.0240	0.3569 2.8016
Kidney	0.0027	0.0000	undef 0.0000
Pancreas	0.0000	0.0055	0.0000 undef
Penis	0.0240	0.0000	undef 0.0000
Prostate	0.0065	0.0021	3.0709 0.3256 undef undef
Uterus-endometrium	0.0000	0.0000 0.0340	0.0000 undef
	0.0000	0.0340	undef undef
Uterus-myometrium	0.0000	0.0000	dider dider
Uterus-general	0.0000		
Breast hyperplasia			
Prostate hyperplasia	0.0118		
Seminal vesicle	0.0000		
Sensory organs	0.0000		
White blood cells		•	
Cervix			

	FETUS % frequency	STANDARDIZED/SUBTE LIBRARIES % frequency	RACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0139 0.0000 0.0000 0.0039 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0036 0.0000 0.0000 0.0000 0.0249 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0000 0.0000 0.0020 0.0000 0.0000 0.0125

Electronic Northern			
	NORMAL	TUMOR	Ratios
	<pre>% frequency</pre>	% frequency	N/T T/N
	_		
Bladder	0.0039	0.0153	0.2542 3.9333
Breast	0.0077	0.0038	2.0416 0.4898
Small intestine	0.0123	0.0000	undef 0.0000
Ovary	0.0060	0.0026	2.3025 0.4343
Endocrine tissue	0.0119	0.0251	0.4755 2.1032
Gastrointestinal	0.0096	0.0139	0.6903 1.4487
Brain	0.0163	0.0246	0.6600 1.5152
Hematopoietic	0.0040	0.0000	undef 0.0000
Skin	0.0110	0.0000	undef 0.0000
Hepatic	0.0095	0.0065	1.4706 0.6800
Heart	0.0180	0.0000	undef 0.0000
Testicles	0.0000 0.0135	0.0117 0.0082	0.0000 undef
Lung	0.0000	0.0153	1.6511 0.6057 0.0000 undef
Stomach-esophagus	0.0069	0.0060	1.1422 0.8755
Muscle-skeleton	0.0000	0.0000	undef undef
Kidney	0.0033	0.0276	0.1197 8.3571
Pancreas	0.0090	0.0267	0.3369 2.9678
Penis	0.0196	0.0128	1.5354 0.6513
Prostate	0.0068	0.0000	undef 0.0000
Uterus-endometrium	0.0000	0.0408	0.0000 undef
	0.0357	0.0000	undef 0.0000
Uterus-myometrium	0.0128		
Uterus-general	0.0089		
Breast hyperplasia	0.0267		
Prostate hyperplasia			•
Seminal vesicle	0.0000	•	
Sensory organs	0.0000		•
White blood cells			
Cervix			

	FETUS % frequency	LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung	0.0000 0.0000 0.0375 0.0039 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle	0.0000 0.0000 0.0000 0.0012 0.0122 0.0000 0.0097
Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0507 0.0124 0.0061 0.0249 0.0000	Testicles Lung Nerves Prostate Sensory Organs	0.0000 0.0082 0.0050 0.0000 0.0000

Electronic Northern f	or SEQ. ID NO.	.: 17	
		rumor	Ratios
	% frequency %	frequency	/ N/T T/N
	1		
Bladder	0.0195	0.0179	
Breast	0.0193	0.0179	1.0896 0.9178
Small intestine	0.0184	0.0165	0.9074 1.1021
Ovary	0.0509	0.0286	1.1122 0.8991 1.7792 0.5620
Endocrine tissue	0.0375	0.0301	
	0.0153	0.0093	1.2453 0.8030 1.6567 0.6036
Gastrointestinal	0.0222	0.0452	0.4909 2.0372
Brain	0.0160	0.0000	undef 0.0000
Hematopoietic	0.0661	0.0000	undef 0.0000
Skin	0.0143	0.0518	0.2757 3.6266
Hepatic	0.0636	0.0000	undef 0.0000
Heart	0.0173	0.0117	1.4759 0.6775
Testicles	0.0177	0.0532	0.3322 3.0104
Lung	0.0483	0.0000	undef 0.0000
Stomach-esophagus	0.0137	0.0960	0.1428 7.0040
Muscle-skeleton	0.0217	0.0685	0.3172 3.1524
Kidney	0.0264	0.0055	4.7864 0.2089
Pancreas	0.0449	0.0000	undef 0.0000
Penis	0.0283	0.0341	0.8317 1.2024
Prostate	0.0135	0.0000	undef 0.0000
Uterus-endometrium	0.0229	0.0679	0.3367 2.9702
Uterus-myometrium	0.0051	0.0000	undef 0.0000
Uterus-general	0.0671		•
Breast hyperplasia	0.0476		
Prostate hyperplasia	0.0356		
Seminal vesicle	0.0333		
Sensory organs	0.0121	•	
White blood cells	0.0532		
Cervix			
CETATY			

+	FETUS % frequency	LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0557 0.0278 0.0688 0.0275 0.0000 0.0000 0.0534 0.0831 0.1014 0.0741 0.0182 0.1247 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus_n	0.0000 0.0000 0.0203 0.0490 0.0169 0.0122 0.0000 0.0259 0.0000 0.0082 0.0050 0.0068 0.0000 0.0167

Electronic Northern	for SEQ. ID NO	.: 18	
	NORMAL '	TUMOR	Ratios
	<pre>% frequency ?</pre>	% frequency	N/T T/N
Bladder	0.0195	0.0179	1.0896 0.9178
Breast	0.0371	0.0357	1.0387 0.9627
Small intestine	0.0245	0.0331	0.7415 1.3487
Ovary	0.0479	0.0390	1.2280 0.8143
Endocrine tissue	0.0358	0.0326	1.0972 0.9114
Gastrointestinal	0.0153	0.0093	1.6567 0.6036
Brain	0.0229	0.0524	0.4376 2.2851
Hematopoietic	0.0147	0.0000	undef 0.0000
Skin	0.0587	0.0000	undef 0.0000
Hepatic	0.0143	0.0518	0.2757 3.6266
Heart	0.0593	0.0000	undef 0.0000
Testicles	0.0115	0.0117	0.9839 1.0163
Lung	0.0145	0.0491	0.2964 3.3743
Stomach-esophagus	0.0290	0.0000	undef 0.0000
Muscle-skeleton	0.0120	0.1140	0.1052 9.5055
Kidney	0.0299 0.0281	0.0616	0.4846 2.0634
Pancreas	0.0281	0.0055	5.0855 0.1966
Penis	0.0449	0.0000	undef 0.0000
Prostate	0.0203	0.0319 0.0000	1.0919 0.9159
Uterus-endometrium	0.0152	0.0679	undef 0.0000
Uterus-myometrium	0.0051	0.0000	0.2245 4.4553 undef 0.0000
Uterus-general	0.0576	0.000	dider 0.0000
Breast hyperplasia	0.0446		
Prostate hyperplasia			
Seminal vesicle	0.0235		
	0.0130		
Sensory organs White blood cells	0.0532		
Cervix	*		

	FETUS % frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0417 0.0333 0.0688 0.0275 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0000 0.0000 0.0253 0.0245 0.0326 0.0122
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0427 0.0867 0.1268 0.0741 0.0182 0.1247 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs	0.0518 0.0000 0.0082 0.0090 0.0137 0.0000 0.0208

Electronic Northern for SEQ. ID NO.: 19				
	NORMAL	TUMOR	Ratios	
	% frequency	% frequency	N/T T/N	
p3 - 33		•		
Bladder		·		
Breast	0.0351	0.0332	1.0561 0.9469	
Small intestine	0.0269	0.0376	0.7146 1.3995	
Ovary	0.0337	0.0992	0.3398 2.9425	
Endocrine tissue	0.0240	0.0312	0.7675 1.3029	
Gastrointestinal	0.0409	0.0075	5.4340 0.1840	
Brain	0.0268	0.0971	0.2761 3.6217	
Hematopoietic	0.0059	0.0113	0.5236 1.9098	
Skin	0.0013	0.2273	0.0059 170.0273	
Hepatic	0.0330	0.0000	undef 0.0000	
Heart	0.0095	0.0388	0.2451 4.0800	
Testicles	0.0223	0.0000	undef 0.0000	
Lung	0.0173 0.0395	0.0351 0.0900	0.4920 2.0326	
Stomach-esophagus	0.0000	0.0900	0.4388 2.2792 0.0000 undef	
Muscle-skeleton	0.0857	0.1920	0.4462 2.2413	
Kidney	0.0136		0.6609 1.5132	
Pancreas	0.0198	0.0221	0.8974 1.1143	
Penis	0.0629	- 0.0800	0.7862 1.2719	
Prostate	0.0087	0.0106	0.8189 1.2211	
Uterus-endometrium	0.0676	0.0000	undef 0.0000	
	0.0686	0.1630	0.4208 2.3761	
Uterus-myometrium	0.0153	0.0000	undef 0.0000	
Uterus-general	0.0735			
Breast hyperplasia	0.0416			
Prostate hyperplasia	0.0000			
Seminal vesicle	0.0235			
Sensory organs	0.0000			
White blood cells	0.0426			
Cervix				

	FETUS % frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate	0.1113 0.0056 0.0000 0.0039 0.0000 0.0000 0.0356 0.0289 0.0000 0.0062 0.1333 0.0249	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate	0.0204 0.0000 0.0000 0.0000 0.0111 0.0244 0.0000 0.0065 0.0000 0.0000 0.0000
Sensory organs	0.0000	Sensory Organs Uterus n	0.0125

Electronic Northern	for SEQ. ID NO NORMAL % frequency	TUMOR	Ratios N/T T/N
Bladder Breast Small intestine Ovary Endocrine tissue Gastrointestinal Brain Hematopoietic Skin Hepatic Heart Testicles Lung Stomach-esophagus Muscle-skeleton Kidney Pancreas Penis Prostate Uterus-endometrium Uterus-myometrium Uterus-general Breast hyperplasia Prostate hyperplasia Prostate hyperplasia Seminal vesicle Sensory organs White blood cells Cervix	0.0858 0.1036 0.1226 0.0958 0.0954 0.0900 0.0658 0.1109 0.0918 0.0333 0.2120 0.0690 0.0696 0.0483 0.0702 0.0652 0.1140 0.1018 0.1090 0.1149 0.0686 0.0458 0.1279 0.1157 0.1059 0.0832 0.1278	0.1048 0.1729 0.1158 0.1197 0.1128 0.1573 0.0924 0.0758 0.0000 0.0412 0.3625 0.1186 0.0460 0.2820 0.1027 0.1270 0.0000 0.1277 0.0000 0.2106 0.1908	0.8185 1.2217 0.5992 1.6690 1.0593 0.9441 0.8009 1.2486 0.8453 1.1830 0.5725 1.7466 0.7120 1.4046 1.4645 0.6828 undef 0.0000 0.3676 2.7200 5.1398 0.1946 0.1904 5.2509 0.5869 1.7040 1.0504 0.9520 0.2491 4.0145 0.6344 1.5762 0.8974 1.1143 undef 0.0000 0.8530 1.1723 undef 0.0000 0.3258 3.0692 0.2402 4.1635
		CMANDABDT 71	en / ciibmb i cmen

	FETUS % frequency	LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate	0.0557 0.2471 0.2189 0.1612 0.0000 0.1560 0.2633 0.1012 0.1014 0.1112	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate	0.0136 0.1595 0.0658 0.0000 0.0216 0.0732 0.0057 0.0259 0.0000 0.1638 0.0211 0.0205 0.0000
Sensory organs	0.3740 0.0126	Sensory Organs Uterus n	0.0333

Electronic Northern			D-4:
	NORMAL	TUMOR	Ratios
	% frequency	% frequence	cy N/T T/N
Bladder	· •		
Breast	0.0273	0.0102	2.6695 0.3746
Small intestine	0.0051	0.0019	2.7221 0.3674
Ovary	0.0276	0.0000	undef 0.0000
Endocrine tissue	0.0180	0.0000	undef 0.0000
Gastrointestinal	0.0136	0.0000	undef 0.0000
Brain	0.0134	0.0046	2.8992 0.3449
Hematopoietic	0.0015	0.0164	0.0900 11.1117
Skin	0.0053	0.0000	undef 0.0000
Hepatic	0.0147	0.0000	undef 0.0000
Heart	0.0048	0.0000	undef 0.0000
Testicles	0.0106 0.0000	0.0412	0.2570 3.8912
	0.0052	0.0117 0.0102	0.0000 undef
Lung	0.0000	0.0000	0.5080 1.9684 undef
Stomach-esophagus	0.0000	0.0060	3.7122 0.2694
Muscle-skeleton	0.0081	0.0000	undef 0.0000
Kidney	0.0033	0.0276	0.1197 8.3571
Pancreas	0.0032	0.0267	0.1123 8.9035
Penis	0.0022	0.0043	0.5118 1.9538
Prostate	0.0135	0.0000	undef 0.0000
Uterus-endometrium	0.0381	0.1019	0.3741 2.6732
Uterus-myometrium	0.0102	0.0000	undef 0.0000
Uterus-general	0.0032		
Breast hyperplasia	0.0000		
Prostate hyperplasia	0.0089		•
Seminal vesicle	0.0000		
Sensory organs	0.0130		
White blood cells	0.0106		
Cervix			

	FETUS % frequency	LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0000 0.0111 0.0000 0.0157 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0204 0.0000 0.0101 0.0490 0.0093 0.0122 0.0057
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0107 0.0000 0.0000 0.0247 0.0000 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs	0.0154 0.0000 0.0010 0.0274 0.0000 0.0000

Electronic Northern			- 1.21
	NORMAL	TUMOR	Ratios
	<pre>% frequency</pre>	% frequency	y N/T T/N
Bladder		_	
Breast	0.0000	0.0102	0.0000 undef
Small intestine	0.0064	0.0207	0.3093 3.2328
Ovary	0.0153	0.0662	0.2317 4.3157
Endocrine tissue	0.0180	0.0156	1.1513 0.8686
Gastrointestinal	0.0204	0.0351	0.5822 1.7176
Brain	0.0153	0.0231	0.6627 1.5090
Hematopoietic	0.0510	0.0257	1.9871 0.5032
Skin	0.0134	0.0000	undef 0.0000
Hepatic	0.0367 . 0.0095	0.0000	undef 0.0000
Heart	0.0466	0.0000 0.0000	undef 0.0000 undef 0.0000
Testicles	. 0.0000	0.0117	0.0000 undef
Lung	0.0249	0.0348	0.7172 1.3943
Stomach-esophagus	0.0193	0.0077	2.5211 0.3967
Muscle-skeleton	0.0377	0.0300	1.2564 0.7959
Kidney	0.0244	0.0274	0.8922 1.1209
Pancreas	0.0083	0.0166	0.4986 2.0057
Penis	0.0329	0.0267	1.2355 0.8094
Prostate	0.0131	0.0149	0.8774 1.1397
Uterus-endometrium	0.0338	0.0000	undef 0.0000
Uterus-myometrium	0.0000	0.0340	0.0000 undef
Uterus-general	0.0153	0.0000	undef 0.0000
Breast hyperplasia	0.0256		
Prostate hyperplasia	0.0238		
Seminal vesicle			
Sensory organs	0.0000		
White blood cells	0.0069 0.0213		-
Cervix	0.0213		
CCTATY			

	FETUS % frequency	STANDARDIZED/SUBTRA LIBRARIES % frequency	CTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate	0.0139 0.0194 0.0125 0.0039 0.0000 0.0000 0.0320 0.0434 0.0000 0.0124 0.0061 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate	0.0000 0.3190 0.0000 0.0000 0.0373 0.0000 0.0114 0.0680 0.0231 0.0000 0.0211 0.0205 0.0205
Sensory organs	0.0251	Sensory Organs Uterus n	0.0458

Electronic Northern for	r SEQ. ID N	0.: 23	× .
]	NORMAL	TUMOR	Ratios
	% frequency	% frequence	cy N/T T/N
		-	
Bladder	0.0741	0.0639	1.1593 0.8626
Breast	0.0691	0.0827	0.8352 1.1973
Small intestine	0.0245	0.0496	0.4943 2.0230
Ovary	0.0689	0.0494	1.3936 0.7176
Endocrine tissue	0.2487	0.5191	0.4791 2.0873
Gastrointestinal	0.0421	0.0879	0.4796 2.0852
Brain	0.1700	0.1037	1.6395 0.6099
Hematopoietic	0.0695	0.0758	0.9175 1.0899
Skin	0.0367	0.4237	0.0866 11.5419
Hepatic	0.0143	0.0388	0.3676 2.7200
Heart	0.0668	0.1375	0.4857 2.0588
Testicles	0.0460	0.0468	0.9839 1.0163
Lung	0.0592	0.0470	1.2590 0.7943
Stomach-esophagus	0.1160	0.0690	1.6807 0.5950
Muscle-skeleton	0.0754 0.0706	0.0960 0.0479	0.7853 1.2735
Kidney	0.0706	0.0552	1.4728 0.6790 1.2265 0.8153
Pancreas	0.0988	0.0352	3.7064 0.2698
Penis	0.0697	0.0660	1.0566 0.9464
Prostate	0.0608	0.0000	undef 0.0000
Uterus-endometrium	0.0152	0.0611	0.2494 4.0097
· 	0.1120	0.0000	undef 0.0000
Uterus-myometrium	0.0480		
Uterus-general	0.0	565	•
Breast hyperplasia	0.0		
Prostate hyperplasia	0.0		•
Seminal vesicle	0.0		
Sensory organs	0.0		
White blood cells	0.0	U J 2	•
Cervix			

•	FETUS frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.1113 0.0805 0.1376 0.1140 0.0000 0.0520 0.0996 0.1951 0.1268 0.1359 0.1030 0.0748 0.0879	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0544 0.0000 0.0203 0.0245 0.0309 0.0610 0.0000 0.0356 0.0077 0.0655 0.0783 0.0547 0.0000 0.0083

Electronic Northern for SEQ. ID NO.: 24 NORMAL TUMOR Ratios % frequency % frequency N/T T/N 0.0546 0.0332 Bladder 1.6428 0.6087 0.0269 0.0320 0.8407 1.1896 Breast 0.0276 0.0000 Small intestine undef 0.0000 0.0479 0.0546 0.8771 1.1401 Ovary 0.0324 0.0176 1.8437 0.5424 Endocrine tissue 0.0211 0.0463 0.4556 2.1950 Gastrointestinal 0.0229 0.0277 0.8266 1.2097 Brain 0.0348 0.0379 0.9175 1.0899 Hematopoietic 0.0184 0.0000 undef 0.0000 Skin 0.0190 undef 0.0000 undef 0.0000 0.0000 Hepatic 0.0477 0.0000 Heart 0.1169 0.0230 0.1968 5.0816 Testicles 0.0156 0.0307 0.5080 1.9684 0.0000 Lung 0.0460 0.0000 undef 0.0120 0.0480 Stomach-esophagus 0.2499 4.0023 0.0163 0.0068 Muscle-skeleton 2.3791 0.4203 0.0182 0.0331 0.5484 1.8234 Kidney 0.0180 0.0800 0.2246 4.4517 Pancreas 0.0174 . 0.0170 1.0236 0.9769 Penis 0.0541 0.0000 undef 0.0000 0.3367 2.9702 Prostate 0.0229 . 0.0679 Uterus-endometrium 0.0102 0.0000 undef 0.0000 Uterus-myometrium 0.0576 Uterus-general 0.0208 Breast hyperplasia 0.0089 Prostate hyperplasia 0.0118 Seminal vesicle 0.0277 Sensory organs White blood cells 0.0213 Cervix

	FETUS % frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0278 0.0639 0.0438 0.0275 0.0000 0.0000 0.0925 0.0831 0.0000 0.0803 0.0667 0.0249 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus_n	0.0000 0.0000 0.0000 0.0000 0.0122 0.0000 0.0077 0.0000 0.0040 0.0000 0.0000

Electronic Northern	NORMAL	D.: 25 TUMOR % frequency	Ratios N/T T/N
	* ITequency	% liequency	N/1 1/N
Bladder	0.0429		
Breast	0.0435	0.0486 0.0771	0.8832 1.1323
Small intestine	0.0307	0.0000	0.5643 1.7720
Ovary	0.0389	0.0000	undef 0.0000
Endocrine tissue	0.0460	0.0301	0.4677 2.1381
Gastrointestinal	0.0287	0.0971	1.5283 0.6543 0.2958 3.3803
Brain	0.0347	0.0534	0.6507 1.5367
Hematopoietic	0.0869	0.0000	undef 0.0000
Skin	0.0551	0.0847	0.6498 1.5389
Hepatic	0.0143	0.0194	0.7353 1.3600
Heart	0.0466	0.0137	3.3923 0.2948
Testicles	0.0173	0.0935	0.1845 5.4203
Lung	0.0457	0.0613	0.7451 1.3421
Stomach-esophagus	0.0290	0.0153	1.8908 0.5289
Muscle-skeleton	0.0223	0.1140	0.1954 5.1183
Kidney	0.0407 0.0314	0.0137	2.9739 0.3363
Pancreas	0.0314	0.0442	0.7105 1.4075
Penis	0.0458	0.0533 0.0617	1.4601 0.6849
Prostate	0.0473	0.0000	0.7412 1.3491
Uterus-endometrium	0.0305	0.1087	undef 0.0000 0.2806 3.5642
Uterus-myometrium	0.0357	0.0954	0.3736 2.6765
Uterus-general	0.0256		0.0.30 2.0703
Breast hyperplasia	0.0803		
Prostate hyperplasia	0.0801		
Seminal vesicle	0.0118		
Sensory organs	0.0494		
White blood cells	0.0426	•	
Cervix			
	•		

	FETUS % frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0417 0.0611 0.0626 0.0708 0.0000 0.0260	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0204 0.1595 0.0101 0.0000 0.0332 0.0000 0.0000
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0534 0.0542 0.0761 0.0988 0.0303 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0077 0.0328 0.0161 0.0068 0.0000 0.0291

Electronic Northern			
	NORMAL	TUMOR	Ratios
	% frequency	<pre>% frequency</pre>	N/T T/N
Bladder	0.0000	0.0000	undef undef
Breast	0.0000	0.0000	undef undef
Small intestine	0.0000	0.0000	undef undef
Ovary	0.0000	0.0000	undef undef
Endocrine tissue	3.0675	0.0000	undef 0.0000
Gastrointestinal	0.0000	0.0000	undef undef
Brain	0.0015	0.0000	undef 0.0000
Hematopoietic	0.0000	0.0000	undef undef
Skin	0.0000	0.0000	undef undef
Hepatic	0.0000	0.0000	undef undef
Heart	0.0000	0.0000	undef undef
Testicles	0.0000	0.0000	undef undef
Lung	0.0000	0.0000	undef undef
Stomach-esophagus	0.0000	0.0000	undef undef
Muscle-skeleton	.0.000	0.0000	undef undef
Kidney	0.0000	0.0000	undef undef
Pancreas	0.0000	0.0000 0.0000	undef undef
Penis	0.0000	0.0000	undef undef
Prostate	0.0000	0.0000	undef undef undef undef
Uterus-endometrium	0.0000	0.0340	0.0000 undef
Uterus-myometrium	0.0000	0.0000	undef undef
Uterus-general	0.0000	3.4444	midel didel
Breast hyperplasia	0.0000		
Prostate hyperplasia	0.0000		
Seminal vesicle	0.0000		
	0.0000	•	
Sensory organs White blood cells	0.0000		
Cervix			
CELAIX			

	FETUS % frequency	LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0000 0.0028 0.0000 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0000 0.0000 0.0000 0.0000 0.0070 0.0000 0.0000
Lung	0.0000	Skin-muscle	0.0000
Suprarenal gland	0.0000	Testicles	0.0000
Kidney	0.0000	Lung	0.0000
Placenta	0.0000	Nerves	0.0000
Prostate	0.0061	Prostate	0.0155
Sensory organs	0.0000 0.0000	Sensory Organs	0.0250

Electronic Northern	NOR	IAL	TUMOR	Ratios	
	% fr	requency	% frequency	Y N/T	T/N
Bladder Breast Small intestine Ovary Endocrine tissue Gastrointestinal Brain Hematopoietic Skin Hepatic Heart Testicles Lung Stomach-esophagus Muscle-skeleton Kidney Pancreas Penis Prostate Uterus-endometrium Uterus-myometrium Uterus-general Breast hyperplasia Prostate hyperplasia Prostate hyperplasia Seminal vesicle Sensory organs White blood cells Cervix	*	0.0039 0.0179 0.0061 0.0300 0.0068 0.0077 0.0015 0.0053 0.0000 0.0032 0.0000 0.0032 0.0000 0.0136 0.0000 0.0136 0.0000 0.0030 0.0051 0.0000 0.0030 0.0051 0.0000 0.0030 0.0089 0.0000 0.0013	0.0230 0.0395 0.0000 0.0130 0.0251 0.0093 0.0092 0.0000 0.0000 0.0194 0.0000 0.0368 0.0230 0.0060 0.0068 0.0276 0.0000 0.0128 0.0000	0.1695 5.89 0.4537 2.20 undef 0.00 2.3025 0.43 0.2717 3.68 0.8283 1.20 0.1600 6.25 undef 0.00 undef unde 0.0000 under undef unde 0.2258 4.428 1.2605 0.79 0.0000 under 1.9826 0.504 0.0000 under undef 0.000 0.5118 1.953 undef 0.000 0.0000 undef undef 0.000 0.0000 undef	42 00 43 05 72 04 00 6 6 00 6 6 00 6 8 8 8 8 8 8 9 9 9 9 9 9 9 9 9 9 9 9 9

	FETUS % frequency	STANDARDIZED/SUBTRA LIBRARIES % frequency	CTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0139 0.0278 0.0125 0.0079 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0000 0.0000 0.0304 0.0245 0.0029
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0320 0.0289 0.0000 0.0185 0.0000 0.0997 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0114 0.0097 0.0000 0.0164 0.0020 0.0068 0.0000

Electronic Northern			.
	NORMAL	TUMOR	Ratios
	% frequency	% frequen	cy N/T T/N
n1 - 11			
Bladder	0.0195	0.0383	0.5085 1.9666
Breast	0.0256	0.0714	0.3582 2.7919
Small intestine	0.0552	0.0331	1.6683 0.5994
Ovary	0.0270	0.0468	0.5756 1.7372
Endocrine tissue	0.0477	0.0451	1.0566 0.9464
Gastrointestinal	0.0326	0.0324	1.0058 0.9942
Brain	0.0503	0.0277	1.8132 0.5515
Hematopoietic	0.0201	0.1894	0.1059 9.4460
Skin	0.0367	0.2542	0.1444 6.9252
Hepatic	0.0476	0.0388	1.2255 0.8160
Heart	0.0699	0.0550	1.2721 0.7861
Testicles	0.0173	0.1403	0.1230 8.1305
Lung	0.0395	0.0818	0.4826 2.0720
Stomach-esophagus	0.0676	0.0613	1.1030 0.9066
Muscle-skeleton	0.0394	0.0300	1.3135 0.7613
Kidney	0.0462 0.0347	0.0616	0.7490 1.3351
Pancreas	0.0347	0.0607	0.5711 1.7510
Penis	0.0327	0.1600 0.0149	0.3182 3.1424
Prostate	0.0000	0.0000	2.1935.0.4559
Uterus-endometrium	0.0000	0.0543	undef undef
	0.0357	0.0000	0.1403 7.1284 undef 0.0000
Uterus-myometrium	0.0192	0.000	dider 0.0000
Uterus-general	0.0565		
Breast hyperplasia			•
Prostate hyperplasia	0.0941		•
Seminal vesicle	0.0390		
Sensory organs	0.0319		
White blood cells			
Cervix			

	FETUS % frequency	LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta	0.0557 0.0194 0.0188 0.0197 0.0000 0.0000 0.0320 0.0325 0.0000 0.0371	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves	0.0340 0.0000 0.0101 0.0000 0.0396 0.0244 0.0000 0.0778 0.0000 0.0000
Prostate Sensory organs	0.0997 0.0000	Prostate Sensory Organs	0.0479
		Uterus_n	0.0291

Electronic Northern fo	r SEQ. ID NO	D.: 29	
	NORMAL	TUMOR	Ratios
,	% frequency	% frequency	N/T T/N
		- 4	_,
Bladder			
Breast	0.0663	0.0358	1.8523 0.5399
Small intestine	0.0512	0.0489 0.0496	1.0470 0.9551 0.7415 1.3487
Ovary	0.0368 0.0779	0.0498	1.0690 0.9354
Endocrine tissue	0.0779	0.0326	1.7242 0.5800
Gastrointestinal	0.0345	0.0370	0.9319 1.0731
Brain	0.0343	0.0524	0.7482 1.3366
Hematopoietic	0.0602	0.0379	1.5880 0.6297
Skin	0.0587	0.0000	undef 0.0000
Hepatic	0.0095	0.0647	0.1471 6.7999
Heart	0.0583	0.0550	1.0601 0.9433
Testicles	0.0173	0.1520	0.1135 8.8080
Lung	0.0208	0.0491	0.4234 2.3620
Stomach-esophagus	0.0387	0.0460	0.8404 1.1900
Muscle-skeleton	0.0360	0.0420	0.8567 1.1673
Kidney	0.0380	0.0616	0.6168 1.6213
Pancreas	0.0314	0.0884	0.3552 2.8150
Penis	0.0689	0.0267	2.5833 0.3871
Prostate	0.0436	0.0490	0.8901 1.1235
• •	0.0541	0.0000	undef 0.0000
Uterus-endometrium	0.0381	0.1155 0.0000	0.3301 3.0296 undef 0.0000
Uterus-myometrium	0.0815 0.0831	0.0000	under 0.0000
Uterus-general	0.0446		
Breast hyperplasia	0.0443		
Prostate hyperplasia	0.0353		
Seminal vesicle	0.0520		
Sensory organs	0.0532		
White blood cells		•	
Cervix			

	FETUS % frequency	STANDARDIZED/SUBTR LIBRARIES % frequency	ACTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0000 0.0389 0.0188 0.0472 0.0000 0.0260 0.0498	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0000 0.1595 0.0203 0.0000 0.0093 0.0366 0.0000
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0614 0.0254 0.0741 0.0364 0.0499	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0.0130 0.0000 0.0164 0.0120 0.0205 0.0000 0.0208

Electronic Northern :			_	
	NORMAL	TUMOR	Ratios	
	% frequency	% frequen	1CY N/T T/N	
			-	
Bladder	0.0078	0.0000	undef 0.0000	
Breast	0.0013	0.0000	undef 0.0000	
Small intestine	0.0092	0.0000	undef 0.0000	
Ovary	0.0000	0.0000	undef undef	
Endocrine tissue	0.0034	0.0000	undef 0.0000	
Gastrointestinal	0.0019	0.0000	undef 0.0000	
Brain	0.0000	0.0000	undef undef	
Hematopoietic	0.0000	0.0000	undef undef	
Skin	0.0000	0.0000	undef undef	
Hepatic .	0.0000	0.0000	undef undef	
Heart	0.0011	0.0000	undef 0.0000	
Testicles	0.0000	0.0000	undef undef	
	0.0010	0.0000	undef 0.0000	
Lung	0.0000	0.0000	undef undef	
Stomach-esophagus	0.0000	0.0000	undef undef	
Muscle-skeleton	0.0000	0.0000	undef undef	
Kidney	0.0000	0.0000	undef undef	
Pancreas	0.0000	0.0000	undef undef	
Penis	0.0109	0.0021	5.1181 0.1954	
Prostate	0.0000 0.0000	0.0000	undef undef	
Uterus-endometrium	0.0102	0.0340	0.0000 undef	
Uterus-myometrium	0.0000	0.0000	undef 0.0000	
Uterus-general	0.0000		·	
Breast hyperplasia	0.0089			
Prostate hyperplasia	.0.0000			
Seminal vesicle	0.0000			
Sensory organs	0.0000		•	
White blood cells				
Cervix				
				

	FETUS % frequency	STANDARDIZED/SUBTRA LIBRARIES % frequency	CTED
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels	0.0000 0.0000 0.0000 0.0000 0.0000	Breast Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000
Lung Suprarenal gland Kidney Placenta Prostate Sensory organs	0.0000 0.0000 0.0000 0.0000 0.0000	Skin-muscle Testicles Lung Nerves Prostate Sensory Organs Uterus n	0000.0 0000.0 0000.0 8000.0 0000.0

Electronic Northern	NORMAL	O.: 31 TUMOR % frequenc	Ratios y N/T T/N
Bladder Breast Small intestine Ovary Endocrine tissue Gastrointestinal Brain Hematopoietic Skin Hepatic Heart Testicles Lung Stomach-esophagus Muscle-skeleton Kidney Pancreas Penis Prostate Uterus-endometrium Uterus-myometrium Uterus-myometrium Uterus-general Breast hyperplasia Prostate hyperplasia Seminal vesicle Sensory organs White blood cells	0.0000 0.0038 0.0031 0.0034 0.0057 0.0111 0.0040 0.0000 0.0048 0.0095 0.0058 0.0042 0.0000 0.0034 0.0163 0.0163 0.0163 0.0017 0.0090 0.0022 0.0135 0.0000 0.0022 0.0135 0.0000 0.0024 0.0000 0.0024	0.0051 0.0019 0.0000 0.0104 0.0100 0.0046 0.0051 0.0000 0.0000 0.0000 0.0000 0.0000 0.0060 0.0066 0.0066 0.0000 0.0021 0.0000 0.0340 0.0000	0.0000 undef 2.0416 0.4898 undef 0.0000 0.2878 3.4745 0.3396 2.9444 1.2425 0.8048 2.1599 0.4630 undef 0.0000 undef undef undef 0.0000 undef 0.0000 undef 0.0000 undef 0.0000 0.4064 2.4605 undef undef 0.5711 1.7510 2.3791 0.4203 undef 0.0000 undef 0.0000 undef 0.0000 undef 0.0000 undef 0.0000 undef undef
Cervix			

Development Breast 0.0000 Gastrointestinal 0.0000 Ovary_n 0.0000 Brain 0.0028 Ovary_t 0.0000 Hematopoietic 0.0125 Endocrine tissue 0.0000 Skin 0.0000 Fetal 0.0000 Hepatic 0.0000 Gastrointestinal 0.0000 Heart-blood vessels 0.0000 Hematopoietic 0.0000 Lung 0.0000 Skin-muscle 0.0000 Suprarenal gland 0.0036 Testicles 0.0000 Kidney 0.0062 Lung 0.0082 Placenta 0.0303 Nerves 0.0000 Prostate 0.0000 Sensory Organs 0.0000		* frequency	LIBRARIES % frequency	,
Uterus n	Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Suprarenal gland Kidney Placenta Prostate	0.0028 0.0125 0.0000 0.0000 0.0000 0.0036 0.0036 0.0254 0.0062 0.0303 0.0000	Ovary_n Ovary_t Endocrine tissue Fetal Gastrointestinal Hematopoietic Skin-muscle Testicles Lung Nerves Prostate	0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0000 0.0082 0.0000

STANDARDIZED/SUBTRACTED

Electronic Northern	NORMAL	52 TUMOR	RATIOS
B lymphoma Bladder Breast Large intestine Small intestine Ovary Endocrine tissue Brain Skin Hepatic Heart Testicles Lung Stomach-esophagus Muscle-skeleton Kidney Pancreas Prostate T lymphoma Uterus White blood cells Hematopoietic Penis Seminal vesicle Sensory organs	% freq. 0.0599 0.0507 0.0326 0.0268 0.0439 0.0564 0.0642 0.0380 0.0330 0.0093 0.0589 0.0482 0.0389 0.0072 0.0240 0.0694 0.0297 0.0443 0.0505 0.0325 0.0325 0.0325 0.0325 0.0322 0.0493	% freq. 0.0543 0.0282 0.0450 0.0313 0.0426 0.0453 0.0530 0.0379 0.0789 0.0508 0.0000 0.0533 0.0499 0.0256 0.0702 0.0289 0.0331 0.0287 0.1643 0.0690 0.0304	N/T T/N 1.1037 0.9060 1.7979 0.5562 0.7240 1.3811 0.8563 1.1679 1.0306 0.9703 1.2443 0.8037 1.2113 0.8255 1.0022 0.9978 0.4190 2.3868 0.1831 5.4614 undef 0.0000 0.9047 1.1054 0.7796 1.2828 0.2833 3.5296 0.3418 2.9256 2.3984 0.4169 0.8974 1.1143 1.5457 0.6470 0.3074 3.2533 0.4715 2.1210 1.1948 0.8370
	FETUS % freq.		
Development Gastrointestinal Brain Hematopoietic Skin Hepatic Heart-blood vessels Lung Adrenal gland Kidney Placenta Prostate Sensory organs	0.0696 0.0750 0.0876 0.0904 0.0000 0.0260 0.0854 0.0867 0.0761 0.0556 0.0606 0.0499		

STANDARDIZED/SUBTRACTED LIBRARIES & frequency

Breast	
Breast t	0.0068
Large Intestine t	0.0000
Ovary n	0.0000
Ovary	0.1595
Ovary_t	0.0101
Endocrine tissue	0.0245
Fetal	0.0243
Gastrointestinal	
Hematopoietic	0.0732
Hematopoietic Skin-muscle	0.0000
	0.0616
Testicles_n	0.0293
Testicles ⁻ t	0.0000
Lungs n	0.0195
Lungs t	-
Nerves	0.0000
	0.0261
Kidney_t	0.0000
Ovary uterus	0.0293
Prostate n	0.0121
Sensory Organs	0.0310
White blood cells	
MILEGO DIGGG CCIIS	0.0000

2.2. Fisher Test

In order to decide whether a partial sequence S of a gene occurs significantly more often or less often in a library for normal tissue than in a library for degenerated tissue, Fisher's exact test, a standard statistical process, is carried out (Hays, W. L., (1991) Statistics, Harcourt Brace College Publishers, Fort Worth).

The null hypothesis reads: The two libraries cannot be distinguished with respect to the frequency of sequences homologous to S. If the null hypothesis can be rejected with high enough certainty, the gene belonging to S is accepted as an advantageous candidate for a cancer gene, and in the next step an attempt is made to achieve lengthening of its sequence.

Example 3

Automatic Lengthening of the Partial Sequence

Automatic lengthening of partial sequence S is completed in three steps:

- Determination of all sequences homologous to S from the total set of available sequences using BLAST
- 2. Assembling these sequences by means of the standard program GAP4 (Bonfield, J. K.; Smith, K. F. and Staden, R. (1995), Nucleic Acids Research 23 4992-4999) (contig formation).
- Computation of a consensus sequence C from the assembled sequences.

Consensus sequence C will generally be longer than initial sequence S. Its electronic Northern Blot will accordingly

deviate from that for S. A repeated Fisher test decides whether the alternative hypothesis of deviation from a uniform expression in the two libraries can be maintained. If this is the case, an attempt is made to lengthen C in the same way as S. This iteration is continued with consensus sequences C_i (i: iteration index) obtained in each case until the alternative hypothesis is rejected (if H_0 Exit; truncation criterion I) or until automatic lengthening is no longer possible (while $C_i > C_{i-1}$; truncation criterion II).

In the case of truncation criterion II, with the consensus sequence present after the last iteration, a complete or roughly complete sequence of a gene which can be related to cancer with high statistical certainty is acquired.

Analogously to the above-described examples, it was possible to find from hysteromyomic tissue the nucleic acid sequences described in Table I.

Furthermore, for the individual nucleic acid sequences, it was possible to determine the peptide sequences (ORF's) that are listed in Table II, in which no peptide can be assigned to a few nucleic acid sequences and more than one peptide can be assigned to some nucleic acid sequences. As already mentioned above, both the determined nucleic acid sequences and the peptide sequences assigned to the nucleic acid sequences are the subject of this invention.

Example 4

Mapping of Nucleic Acid Sequences on the Human/Genom

Human genes were mapped using the Stanford G3 Hybrid Panel Stewart et al., 1997), which is marketed by Research Genetics, Huntsville, Alabama. This panel consists of 83 different genomic DNAs of human-hamster hybrid cell lines and allows resolution of 500 kilobases. The hybrid cell lines were obtained by fusion of irradiated diploid human cells with cells of the Chinese hamster. The retention pattern of the human chromosome fragments is determined by means of gene-specific primers in a polymerase chain reaction and is analyzed using software available from the Stanford RH server (http://www.stanford.edu/RH/rhserver_form2. html). This program determines the STS marker that is nearest to the desired gene. The corresponding cytogenetic band was determined using the "Mapview" program of the Genome Database (GDB), (http://gdbwww.dkfz-heidelberg.de).

In addition to mapping of genes on the human chromosome set by various experimental methods, it is possible to determine the location of genes on this by biocomputer methods. To do this, the known program e-PCR was used (Schuler GD (1998) Electronic PCR: Bridging the Gap between Genome Mapping and Genome Sequencing. Trends Biotechnol 16: 456-459, Schuler GD (1997). Sequence Mapping by Electronic PCR. Genome Res. 7: 541-550). The database used here no longer corresponds to the one cited in the literature, but is a further development which includes data from the public database RHdb (http://www.ebi.ac.uk/RHdb/-index.html). Analogously to the mapping by the hybrid panels,

the results were evaluated with the above-mentioned software and the software of the Whitehead Institute (http://carbon.wi.mit.edu:8000/cgi-bin/contig/rhmapper.pl).

CONTE

```
TABLE I
```

```
Col. 1 - Sequence ID No.:
Col. 2 - Expression in hysteromyomic/tissue:
Col. 3 - Function
Col. 4 - Modules
Col. 5 - Length of the applied sequence in bases
Col. 6 - Cytogenetic localizatión
Col. 7 - Next marker
[Key to Table I:]
[Col. 2:]
           erhöht = elevated
[Col. 3:]
     [Seq. ID No.: 14, 1/5, 17, 30, 31] unbekannt = unknown
     [Seq. ID No.: 16]
                         Homolog zu Homo sapiens... = homologous
                          to homo sapiens...
     [Seq. ID No.: 18] Humanes Homolog zu ... = human homolog to
                         Verlängerung von Seq. ID. 14 =
  Lengthening of Seq. ID. 14
     [Seq. ID No.: 52]
```

TABLE I

				D5S1730		D2S387	0000	. 08/6-IVV							WI-4204					D12S351												WI-4204
Lokalisation				5q11.2-q13.1	1p36.11-p36.13	2p23.3	5q32-q33.1	2p22.3-p22.1			7-40 0 043	/p12.2-p13			14p11.2-14p11.1					12q21.31-12q21.33				20q13.32-q13.33					1p32.3-p34.3			14011 2 14011 1
Lange der angemeldeten Sequenz in	Basen	779	2310	854	1112				999	1775	506	2191	1769	1026	676	1254	537	823	1682	1548	844	862	546	1591	441	1131	1071	968	1050	581	264	111
Module an			"abhydro- lase"				"kazal"	ູ,ແມ,	3x "TIM"		"Thymosin"	"IGFBP", "Ihyro-	Alabania i	"lipocalin"	"rrm"			1		5x "LRR"		"PMP22"	-	"G-alpha",	ູ,ແມ	"HMG14_17"	"hormone"	3x "PTN MK"	"AhpC-TSA"			3
Funktion		Human mRNA for ornithine decarboxylase antizyme		Human cocaine and amphetamine regulated transcript	CART (nCART)	Human mRNA for KIAA0108 gene		Homo sapiens splicing factor, arginine/serine-rich 7	riosephospi			Human growth hormone-dependent insulin-like growth factor-binding protein mRNA		Human H19	Iniai leili	unbekand	Homolog zu Homo sapiens mRNA for putalively	prenylated protein	Unbekannt	_!^	46	associated prolein	Human YMP		Human huRNP core protein A1	Human HMG-17 gene for non-histone chromosomal	H. sapiens mRNA for prolactin (clone PRL 205)	Human mRNA for neurlle outgrowth-promoting protein.	H saplens mRNA for proliferation-associated gene (pag)		unbekannt	unbekannt
Expression im Uterus-	муотремере	erhöhl	erhöht	erhöhl		erhöhl	erhöhl	erhöht	/erhöht	- Prhibit	erhähl	erhöhl		erhöht	erhöhl	ernoni	erhöht		erhöhl	ernoni	erhöhl		erhäht	erhöhl	ochähl	erhöht	erhöht	erhöht	prhöht	erhöhl	erhöht	erhöht
Sequenz ID No.:		-	2	3		4	2 4	-	/ -	0	٦	=		12	13	4	<u>c</u> 9		-	B	20		21	22		25	26	12	ōC	200	8	31

References to the modulés:

Pfam: Protein families database of alignments and HMMs (pfam@sanger.ac.uk)

PROSITE: The PROSITE database, its status in 1999. Nucleic Acids Res. 27: 215-219 (http://www.expasy.ch/sprot/prosite.html)

للمرخة

TABLE II

DNA Sequ	ences	Peptide Sequences	(ORF's)
Seq. ID.	No.	Seq. ID. No.	
4.4		32	
14			
		33	
•		34	
15		35	
		36	
<u> </u>		37	
16	·	38	,
		39	
16 17 18		40	
		41	
.— Ti		42	
[N 18		43	
된 	• (4)	44	
		45	
30		46	
		47	
31		48	
		49	,
•		50	
		51	·
	·		
52		53	•
		54	
		55	

The inventive nucleic acid sequences Seq. ID No. 1 to Seq. ID No. 31 of the determined candidate genes and the determined amino acid sequences Seq. ID No. 32 to Seq. ID No. 51 are described in the following sequence protocol.

Sequence Protocol

- (1) GENERAL INFORMATION:
 - (i) APPLICANT:
 - (A) NAME: metaGen Gesellschaft für Genomforschung mbH
 - (B) STREET: Ihnestrasse 63
 - (C) CITY: Berlin
 - (E) COUNTRY: Germany
 - (F) POSTAL CODE (ZIP): D-14195
 - (G) TELEPHONE: (030)-8413 1673
 - (H) FAX: (030)-8413 1674
 - (ii) TITLE OF INVENTION: Human Nucleic Acid Sequences from Hysteromyomic Tissue
 - (iii) Number of sequences: 55
 - (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: Patentin release #1.0, version #1.25 (EPO)

- (2) INFORMATION ON SEQ ID NO. 1:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 779 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual
 - ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

agegageage ggeggeggeg eggagagaeg eageggaggt tttcctggtt teggacecca 60 geggeeggat ggtgaaatee teeetgeage ggateeteaa tageeactge ttegeeagag120 agaaggaagg ggataaaccc agcgccacca tccacgccag ccgcaccatg ccgctcctaa180 gcctgcacag ccgcggcggc agcagcagtg agagttccag ggtctccctc cactgctgta240 gtaacceggg teeggggeet eggtggtget cetgatgeee eteacceace eetgaagate300 ccaggtgggc gagggaatag tcaaagggac cacaatcttt cagctaactt attctactcc360 gatgategge tgaatgtaac agaggaacta acgtecaacg acaagacgag gattetcaac420 gtccagtcca ggctcacaga cgccaaacgc attaactggc gaacagtgct gagtggcggc480 actgetetae ategagatee egggeggege getgeeegag gggageaagg acagetttge540 agttctcctg ggagttcgct gaggagcagc tgcgaggccg accatgtctt aatttgcttc600 cacaagaacc cogaggacag agoogcottg otocgaacct toagottttt cgggcttgag660 attitigagae eggggeatee cettigitee ecaagagace egaegetige ticatgggee720 tacaagtttc gagagagagt ctttggggag aggaagaagg attaggggcc gcgtcgggt 779

- (2) INFORMATION ON SEQ ID NO. 2:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 2310 base pairs
 - (B) TYPE: Nucleic acid (C) STRAND: individual

 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO

- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

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gttctccgaa acatggagtc ctgtaggcaa ggtcttacct gaatcaggat gagggagtgg
tgggtccagg tggggctgct ggccgtgccc ctgcttgctg cgtacctgca catcccaccc 120
ceteageget eccetgeest teacteatgg aagtetteag geaagttitt cacttacaag 180
ggactgcgta tcttctacca agactctgtg ggtgtggttg gaagtccaga gatagttgtg 240
cttttacacg gttttccaac atccagctac gactggtaca agatttggga aggtctgacc 300
ttgaggtttc atcgggtgat tgcccttgat ttcttaggct ttggcttcag tgacaaaccg 360
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caggagette tetacaggta caagcagaat egatetggte ggettaceat aaagagtete 540
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gacatgtggg cagggatccg caacaatgac gggaacttag tcattgacag tctcttacag 780
tacatcaatc agaggaagaa gttcagaagg cgctggmtgg gagctcttgc ctctgtaact 840
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gagetgtaca ggaaaacget geogeggtee acagtgtaga ttetggatga ceacattage 960
cactatccac agetagagga teccatggge ttettgaatg catatatggg etteateaac1020
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acaaagtcca ctttactcaa attggtgaac agtgtatagg aagaagccag caggagctct1200
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quandacett cttttecata anagactttt tttaacactt tegacttctc teanatattt1380
aqaaqtqcta atttctqqcc cacccccaac aqqaattcta taqtaaqqqq qaqqaqaqq1440
gggctcctt ccctctcctc gaatgacgtt atgggcacat gccttttaaa agttctttaa1500
qcaacacaga gctgagtcct ctttgtcata cctttggatt tagtgtttca tcagctgttt1560
ttagttataa acattttgtt aaaatagata ttggtttaaa tgatacagta ttttaggtat1620
gatttaagac tatgatttac ctatacatta tatatatttt ataaagatac taaaccagca1680
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agacaggaaa gttccagaaa ctttaagaac aaactctgaa agacctatga gcaaatggtg2100
ctgaatactt tttttttaaa gccacatttc attgtcttag tcaaagcagg attattaagt2160
gattatttaa aattogtitt titaaattag caacitcaag tataacaaci tigaaacigg2220
gcttttgaag tctatgtgtg gggggggggt
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- (2) INFORMATION ON SEQ ID NO. 3:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 854 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual

ESTs by assembling and editing

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

ctgcacggg gctcgggctc actataaaag gtgggagcgc gtggtgcccc agcaacgacg 60 agtttcagaa cgatggagag ctcccgcgtg aggctgctgc ccctcctggg cgccgccctg120 ctgctgatgc tacctctgtt gggtacccgt gcccaggagg acgccgagct ccaagcccga180 gccctggaca tcttactctgc cgtggatgat gcctcccacq agaaggagct gatcgaagcg240 ctgcaagaag tcttgaagaa gctcaagagt aaacgtgttc ccatctatga gaagaagtat300 gggaagctgt gtgactgtcc ccgagggaacc tcctgcaatt ccttcctcct gaagtgctta420 tgaaggggcg tccattctcc tccataacatc cccatcctcc tactttcccc agagggaccac480 accttcctcc ctggagtttg gcttaagcaa cagaataaag tttttatttc ctctgaaggg540 aaagggctct tccatagat gtgaagaacac tccttgt gaagagtatt ctcttattta tttgaaaagggaaaccactttgt gtaacctttgt gtaaagaag gaagctttgt ttgaaaaatt tcttatttta tttgtctgac660 aaactcttgt gtacctttgt gtaaagaag gaagctttgt ttgaaaaattg tatttttgta720 tgtggcatgg cagaatgaaa attagatcta gaagcatgta caaattatac attagatg40 gttttaata attg

- (2) INFORMATION ON SEQ ID NO. 4:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1112 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

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cotgootott tottgooatg agagotgoot acctottoot gotattootg cotgoaggot 180
tgctggctca gggccagtat gacctggacc cgctgccgcc gttccctgac cacgtccagt 240
acacccacta tagcgaccag atcgacaacc cagactacta tgattatcaa gaggtgactc 300
ctoggocoto ogaggaacag ttocagttoo agtocoagoa goaagtocaa caggaagtoa 360
teccaquece aaceccagaa ecaggaaatg cagagetgga geccacagag eetgggeete 420
ttgactgccg tgaggaacag tacccgtgca cccgcctcta ctccatacac aggccttgca 480
aacagtqtct caacgaggtc tgcttctaca gcctccgccg tgtgtacgtc attaacaagg 540
agatetqtqt teqtacagtq tqtqcccatq agqaqetect ceqagetqae etetqteqqq 600
acaagttete caaatgtgge gtgatggeea geageggeet gtgeeaatee gtggeggeet 660
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caaaattaaa aacaagtttt tacaaaaaaa aa
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- (2) INFORMATION ON SEQ ID NO. 5:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1051 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:

- (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

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tgggttttgg aggccgtgaa acagccgttt gagtttggct gcgggtggag aacgtttgtc 120
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cgatcatcct ggggacctgg tacatggtag taaacctatt gatggcaatt ttgctgactg 300
tggaagtgac tcatccaaac tccatgccag ctgtcaacat tcagtatgaa gtcatcggta 360
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agtacgtttt gccaacctat gaaatggccg tgaaaatgcc tgaaaaagaa ccaccacctc 840
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catctggtta aacggccttt ttgggttttt t
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- (2) INFORMATION ON SEQ ID NO. 6:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1516 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual

ESTs by assembling and editing

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

gttgtcctca tccctctcat acagggtgac caggacgttc ttgagccagt cccgcatgcg 60 cagggggaag aagatccatg agaaggagaa gcgcctggag gcaggagacc accccgtgga 120 gctgctggcc cgggacttcg agaagaacta taacatgtac atcttccctg tacactggca 180 gttcggccag ctggaccage accecattga cgggtacete teceacaceg agetggetee 240 actgcgtgct cccctcatcc ccatggagca ttgcaccacc cgctttttcg agacctgtga 300 cctggacaat gacaagtaca tcgccctgga tgagtgggcc ggctgcttcg gcatcaagca 360 gaaggatato gacaaggato tigitgatota aatocactoo ticcacagta coggatioto 420 totttaacco toccottogt gittococca atqittaaaa tqittiggatg gittiqtiqti 480 ctgcctggag acaaggtgct aacatagatt taagtgaata cattaacggt gctaaaaatg 540 adaattctaa cccaagaaca tgacattctt agctgtaact taactattaa ggccttttcc 600 acacgcatta atagtcccat ttttctcttg ccatttgtag ctttgcccat tgtcttattg 660 ggcacatggg gtggacacgg atctgctggg ctctgcctta aacacacatt gcagcttcaa 720 cttttctctt tagtgttctg tttgaaacta atacttaccg agtcagactt tgtgttcatt 780 teattteagg gtettggetg cetgtggget teeceaggtg geetggaggt gggeaaaggg 840 aagtaacaga cacacgatgt tgtcaaggat ggttttggga ctagaggctc agtggtggga 900 gagatecetg cagaacecae caaceagaae gtggtttgee tgaggetgta aetgagagaa 960 agattctggg gctgtgttat gaaaatatag acattctcac ataagcccag ttcatcacca1020 tttcctcctt tacctttcag tgcagtttct tttcacatta ggctgttggt tcaaadtttt1080 gggagcacgg actgtcagtt ctctgggaag tggtcagcgc atcctgcagg gcttctcctc1140 ctctgtcttt tggagaacca gggctcttct caggggctct agggactgcc aggctgtttc1200 agccaggaag gccaaaatca agagtgagat gtagaaagtt gtaaaataga aaaagtggag1260 ttggtgaatc.ggttgttctt tcctcacatt tggatgattg tcataaggtt tttagcatgt1320 tecteetttt etecacette ecetttttte ecceaagaat acagagaaaa eteaaagtta1380 atggggaggg tcggatccta caggcctgag aatcggtcaa ctccaagcat ttcatggaaa1440 aggeggette ctaattaate ctacaaacce ccacccagga tggtgagggg tttcaccaat1500 tcctccaaaa ataaaa

(2) INFORMATION ON SEQ ID NO. 7:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 2367 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

gctgctggt aaagtcgtgt gtcagctttt gcgtcact agcctgggac 60 gctgcttgtt aaagaccga gcacgcgggt ctgtcatcat gtcggtac ggggggtcg 120 gagagagaaac caagttattt gttggtaacc tgggaactgg cgctggcaaa ggggaggttg 180 aaagggcttt caagttatt gttggtaacc tgggaactgg cgctggcaaa gggagatgtag 180 atggaaagg gatttgtgg tacccttaa gagatgcaga aatcctccag 240 atggaaagg gatttgtgg tcccgacg gtcctttga tccaaaagga atgcctcaga 360 gatcacgttt tgatagacca cctgccgac gtccctttga tccaaaagga agatgcatgt 420 acgaggacaa ggdaactat tatgcttatg attgctactg tacaaagaga agatgctag 420 gcaggtcacg gtctagaaca cattctccga ccaagagaag gcgatactc cgctcacgac gtccagacgacgacgacgacgacgacgacgacgacgacgacg							
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gatcaegttt tgatagaca cattgoccgac gtocctttga tocaaatgat agatgctag 420 agatgtggoga gtotagatca cattgotcat catagacag gtotacagaca gtotagatca cattcoccga cagagagaag gogatacact cgccaagcag 540 gcaggagacag gtotagatca cattcoccaga agatcaaga totacagaa gogatacacaga totatoctc 600 acgattattoca aagatcaaga tocaacgaa gatctaggtc tggtcatat aaaggacga 720 gcagcogatc aaagtccaga totccatcc caaaaaagaag toctattoca caagatcaaga toctcoccaga agatcaaga totccagaa gtotattoca acagatcaga toctcoccaga agatcaaga toctcoccaga gtotattoca caagatcaga toctcoccaga agatcaaga toctcoccaga tocacagaa 720 gcagcogatc aaagtcact gaaagaatga totccatocc caaaaaagag tocttocca tocaggaagtc 780 coccagaagaataat caaccatca atcaaaaagag atctgatca cattgatcaa toctgaagac 780 aaagtaaata tocaccatca atcaaaaagag atctgatca cattgataaa toctgaagac 780 aaagtaaata tocaccatca atcaaaaagag atctgatca cattgataaa toctgaagac 960 aagataaata tocaccatca atcaaaaagag atctgatca cattgataaa tocacagag 960 aagataaata tocaccatca atcaaaaagag atctgatca cattgaaaa gtottocca acattataaa acttottaa acattottaa acattottaa acattottaa acattottaa acattataa aggatctaga acataataat tgttaacca ttgataaata accttottaaa ttgatacaa ggactttgaa acattataaa tgttaaccat atcaataataa accttottaaa acattottaa acattataa tgttaccaa acattataaa tgttaccaa acattataaa tgttaccaa acattataaa tgttaccaa acattataaa tgttaccaa acattataaa accttottaa acattataaa accttottaa acattataaa tgttaccaa acattataaa accttottaaa accttottaaa acattataaa accttotaaa acattataaa accttotaaaccaa acattataaa accttottaaa acattataaa accttotaaaccaa acattataaa accttotaaaccaa accttotaaaccaa accttotaaaccaa accttotaaaccaa accttotaaaccaa accttotaaaccaa accttotaaaccaa accttataa accaaaaaaa accttotaaaccaa accttotaaaccaa accttotaaaccaa accttotaaaccaacaa accttotaaacaacaa accttotaaacaacaa accaacaacaa tocaacaacaa accaacaacaa accaacaacaa accaacaa	gatttgcctt	: tgtggaattc	: gaagatccta	gagatgcaga	agatgcagta	cgaggactgg	300
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- (2) INFORMATION ON SEQ ID NO. 8:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 568 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual

ESTs by assembling and editing

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

caccagaçes gageastaga especial especial

- (2) INFORMATION ON SEQ ID NO. 9:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1775 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN

(C) ORGAN:

(vii) OTHER ORIGIN:

(A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

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 tetecetece ettettgtga gatttttttg atetteaget acattttegg etttgtgaga 180
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gattgctggc caggitttag atattaacct ggctgcagag ccaaaagtga accgaggaaa 480
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(2) INFORMATION ON SEQ ID NO. 10:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 509 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:

- (A) ORGANISM: HUMAN
- (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

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- (2) INFORMATION ON SEQ ID NO. 11:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 2191 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

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catcaagaaa gggcatgcta aagacagcca gcgctacaaa gttgactacg agtctcagag 360
cacagatace cagaactict cotcogagic caagegggag acagaatatg giocotgoog 420
tagagaaatg gaagacacac tgaatcacct gaagttcctc aatgtgctga gtcccagggg 480
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tgccgcaagg ttaatgtgga gctcaaatat gccttatttt gcacaaaaga ctgccaagga 720
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(2) INFORMATION ON SEQ ID NO. 12:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1769 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual

ESTs by assembling and editing

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

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gcggggcgga gtgaatgagc tctcaggagg gaggatggtg caggcagggg tgaggagcgc 600
agggggcggc gagcgggagg cactggcctc cagagcccgt ggccaaggcg ggcctcgcgg 660
geggegacgg ageegggate ggtgeeteag egtteggget ggagaegagg eeaggtetee 720
agetggggtg gacgtgccca ccagetgccg aaggcaagac gccaggtccg gtggacgtga 780
caagcaggac atgacatggt ccggtgtgac ggcgaggaca gaggaggcgc gtccggcctt 840
cotgaacaco ttaggotggt ggggotgcgg caagaagcgg gtotgtttot ttacttooto 900
cacggagteg geacactatg getgecetet gggeteecag aacceacaac atgaaagaaa 960
tggtgctacc cagetcaage etgggeettt gaateeggae acaaaaceet etagettggal020
aatgaatatg ctgcacttta caaccactgc actacctgac tcaggaatcg gctctggaag1080
grgaagctag aggaaccaga corcarcago coaacarcaa agacaccaro ggaacagcag1140
egecegeage acceaececg caceggegae tecatettea tggccaecec etgeggeggal200
tocccaccac officerette fictititica footforgie tottigitic fgagetffee1320
tgtctttcct tttttctgag agattcaaag cstccacgac tctgtttccc ccgtccttc1380
tgaatttaat tigcactaag tcattigcac tggttggagt tgtggagacg gccttgagtc1440
teagtacgag tgtgegtgag tgtgageeac ettggeaagt geetgtgeag ggeeeggeeg1500
contocator gagoragata arragadede adortatata codadacete accordecet1560
egectagtet ggaageteeg acegacatea eggageagee tteaageatt ccattacgee1620
ccatctcgct ctgtgcccct ccccaccagg gcttcagcag gagccctgga ctcatcal680
pasasasas asasasasa asasasas
```

(2) INFORMATION ON SEQ ID NO. 13:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1026 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual
 - ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

```
aaaagctgtc cgcgcgggga gcccagggcc agctttgggg ttgtccctgg acttgtcttg 60
gttccagaac ctgacgaccc ggcgacggcg acgtctcttt tgactaaaag acagtgtcca 120
gtgctccage ctaggagtet acggggaceg cetecegege egecaceatg eccaacttet 180
ctggcaactg gaaaatcatc cgatcggaaa acttcgagga attgctcaaa gtgctggggg 240
tgaatgtgat gctgaggaag attgctgtgg ctgcagcgtc caagccagca gtggagatca 300
aacaggaggg agacactttc tacatcaaaa cctccaccac cgtgcgcacc acagagatta 360
acttcaaggt tggggaggag tttgaggagc agactgtgga tgggaggccc tgtaagagcc 420
tggtgaaatg ggagagtgag aataaaatgg tctgtgagca gaagctcctg aagggagagg 480
gccccaagac ctcgtggacc agagaactga ccaacgatgg ggaactgatc ctgaccatga 540
eggeggatga egttgtgtge accagggtet acgteegaga gtgagtggee acaggtagaa 600
cogoggooga agodoaccao tggodatgot cacogodotg ottoactgod cootcogtod 660
caccccctcc ttctaggata gcgctcccct taccccagtc acttctgggg gtcactggga 720
tgeetettge agggtettge trrettrgae etettetete eteccetaca ecaacaaaga 780
ggaatggctg caagagccca gatcacccat teegggttea eteecegeet eeecaagtca 840
gcagtectag ecceaaacca geceagagea gggtetetet aaaggggaet tgagggeetg 900
ageaggaaag actggeeete tagettetae eetttgteee tgtageetat acagtttaga 960
aaaaaa
```

(2) INFORMATION ON SEQ ID NO. 14:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 676 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

ggccattttg tqaaqaqacq aaqactqagc ggttgtggcc gcgttgccga cctccaqcaq 60 cagtcggctt ctctacqcaq aacccgggaq tagqaqactc aqaatcqaat ctcttctccc120 tccccttctt gggcaqcaaq gcgaacccca tccctactca ctggaqctca gctttgattt180 ttaacctcc ttccccaccc ttccaqaaca cacaacattcc attccaaaac tgattttata240 aaqacatttt aaacataatq atgcaacttg gtgtgcacta cagcaaatqt acaggtgttt300 ttttttaat tgtttccaaa accgggacct ggatttaaqa tgtaattttt aaaatttcta360 tttctattt ttcggcagca gttgggttaq aggaggagga gccttttagc ctccccaqaaa420 ctgacctctc tacttcctcq tgtattttta agattgattq atgatgtgga aaqggctttq480 cttgtctgct actgaaaact ttatccttqc ggtttttgtg gaactgcgtt tggaaaqaqa540 aaaqaaatqa actttactga cttgacattt tgcacctccc ggttttcqaa tctgggcaat600 ttagcattt

- (2) INFORMATION ON SEQ ID NO. 15:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1254 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

eggetegage agetegageg geteaaacae eteattigae etigeeaget gaeetteaaa 60 ccctgcattt gaaccgacca acattaagtc cagagagtaa acttgaatgg aataacgaca 120 ttccagaagt taatcatttg aattctgaac actggagaaa aaccgaaaaa tggacggggc 180 atgaagagac taatcatctg gaaaccgatt tcagtggcga tggcatgaca gagctagagc 240 tegggeceag ecceaggetg cageceatte geaggeacee gaaagaactt ecceagtatg 300 gragetcorg aaaggacatt tergaagate aactatatet teergreeat teegarggaa 360 tttcagttca tcagatgttc accatggcca ccgcagaaca ccgaagtaat tccagcatag 420 cggggaagat gttgaccaag gtggagaaga atcacgaaaa ggagaagtca cagcacctag 480 aaggcagege etectettea etetectetg attagatgaa actgttacet taccetaaac 540 acagtatttc tttttaactt ttttatttgt aaactaataa aggtaatcac agccaccaac 600 attecaaget accetgggta cetttgtgca gtagaageta gtgageatgt gageaagegg 660 tgtgcacacg gagactcatc gttataattt actatctgcc aagagtagaa agaaaggctg 720 gggatatttg ggttggcttg gttttgattt tttgcttgtt tgtttgtttt gtactaaaac 780 agtattatet titigaatate giagggacat aagtatatae atgitateea atcaagatgg 840 ctagaatggt gcctttctga gtgtctaaaa cttgacaccc ctggtaaatc tttcaacaca 900 cttccactgc ctgcgtaatg aagtittgat tcatttttaa ccactggaat ttttcaatgc 960 egicattite agitagatga tittgeacti tgagattaaa atgeeatgie tattigatta1020 gtottatttt tttattttta caggottato agtotcactg ttggctgtca ttgtgacaaa1080 gtcaaataaa cccccaagga cgacacacag tatggatcac atattgtttg acattaagct1140 tttgccagaa aatgttgcat gtgttttacc tcgacttgct aaaatcgatt agcagaaagg1200 catggctaat aatgttggtg gtgaaaataa ataaataagt aaacaaaaag aaaa

(2) INFORMATION ON SEQ ID NO. 16:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 537 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual
 - ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEO ID NO: 16:

ggecegggce cccacctcg acatgcgctt ccggcgacgc cttagcgctg accccacgc 60 aacccagcga aactccgcgg aggcgcggg cacgatggac ggtcgggtgc agctgatgaa120 ggccctcctg gccggccc tccggcccqc ggcgcgtcgc tggaggaacc cgattccctt180 tcccqagacg tttgacggag ataccgaccg actcccggag ttcatcgtgc agacgtgctc240 ctacatgttc gtggacgaga acacgttctc caacgacgc ctgaaggtga cgttcctcat300 cacccgcctc acggggccag ccctgcagtg ggtgatcccc tacatcagga aggagagccc360 cctgctcaat gattaccggg gctttctggc cgagatgaag cgagtctttg gatggagga420 ggacgaggac ttctaggcc ggagaccctc gggcctgggg gcgggtgctc tgggaagagt480 tcgctgtgcc agtggccac gctagggtc ccacaggcg cctcccagg gaatgct 537

(2) INFORMATION ON SEQ ID NO. 17:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 823 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

```
tagactgaac aggagggga gtectgggta gegegeeggt etaaategtt acttggegga 60

aagttecat gagtetttge cagegteec eteettttgt gaggattggg atattecegae120

teettaaggg eetggege ataaaggtgtg acetttteat teeegttgtt atggagggee180

acatetgeea gageetggag tetgegaagg eegggaeeeg gtteeeegge ecacagtggg240

ggtgtgeaaa eegggeeagg ageetggea ageetgeeat gaggttttga gaacagagtg etgtttaga360

getggeagea geateteage eeaaggaaag gttatattee eagaggatgt eagteeeaag420

gaacagtage tgeeateagt ttggattetg aaaaetaaet ggeateaaea etgggtgtag480

aaacatgett geettatgta teagaggaea tgeteageag ateeaagaga tatatttgge540

aaacttttet agaaaaggea eattgggtat eagteeagga gtgtggtgg660

tittttttt tttttttga gaeagtettg etgtattgee eaggetgga gtgtggtgge660

acaateaeag eteattgeat eeteaateae eeagggeeta ageaateete eeacettgta720

getgggaeta eageteaeag eacaeeggge taaaaatttt ttttgttgag aeggttttee780

tatgttgeee gggtggttt eagggteegg ggtteagatg gte 823
```

- (2) INFORMATION ON SEQ ID NO. 18:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1082 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:

```
gggcgcacat aaggtgtgac cttttcattc ccgttgttat ggagggccac atctgccaga
geotygagte tgegaaggee gggaeeeggt teeeeggeee acagtggggg tgtgcaaace 120
cgagagaact ggtcgctgaa acctctacaa cttagttgac cgtaactgcc agagccctgc 180
cetgaattee tgteettact ecetetttaa gattgegtae eeactgeaga gtgetgaaga 240
cggggtagcc acgaggttgc aaattcgtga agaatcagca tcatgtttgg cagctgagta 300
ttggagccag gagcctgcca tgaggttttg agaacagagt gctgttttag agctggcagc 360
agcatotoag cocaagagaa ggttatatto coagaggatg toagtocoaa ggaccagtag 420
ctgccatcag tttggattct gaaaactaac tggcatcaac actgggtgta gaaacatgct 480
tgccttatgt atcagaggac atgctcagca gatccaagag atatatttgg caactttttc 540
tagaaaaggc acattgggta tcattcatta cattcttgag tttttttggg ttttttttt 600
ttttttttga gacagtcttg ctgtattgcc caggctggag tgtggtggca caatcacagc 660
teattgeate eteaateace caggeetaag caateeteec acettgtage tgggactaca 720
geteacagea cacetggeta aaattttttt tttgttgaga eggattetet atgttgeeca 780
ggctggtctc aggctcctgg gctcagatgg tcctcctgcc tcagcttcca aaggcacagg 840
ccaagtigta gettigieee tigecateat geccaacaag aggitetata cettitaatg 900
aattgacttt cataaattgg ttatgttggt gggcaagttc tttaagctgg aaattgtaaa 960
tteeteetga aatgittitt eatgeagtta eeatgaacta atactacaat aaaggatggt1020
1082
aa
```

(2) INFORMATION ON SEQ ID NO. 19:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1548 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:

```
cccattccat agggaatgag ctgggctgtc ctttctcccc acgttcacct gcacttcgtt 60
agagagcagt gttcacatgc cacaccacaa gatccccaca atgacataac tccattcaga 120
gactggcgtg actgggctgg gtctccccac ccccccttc agctcttgta tcactcagaa 180
tetggeagee agricegies tgacagagit cacageatat artiggiggat tettgiesat 240
agtgcatctg ctttaagaat taacgaaagc agtgtcaaga cagtaaggat tcaaaccatt 300
tgccaaaaat gagtctaagt gcatttactc tcttcctggc attgattggt ggtaccagtg 360
gocagtacta tgattatgat tttoccotat caatttatgg goaatcatca ccaaactgtg 420
caccagaatg taactgccct gaaagctacc caagtgccat gtactgtgat gagctgaaat 480
tgaaaagtgt accaatggtg cctcctggaa tcaagtatct ttaccttagg aataaccaga 540
ttgaccatat tgatgaaaag gcctttgaga atgtaactga tctgcagtgg ctcattctag 600
atcacaacct totagaaaac tocaagataa aagggagagt tttototaaa ttgaaacaac 660
tgaagaagct gcatataaac cacaacaacc tgacagagtc tgtgggccca cttcccaaat 720
ctctggagga tctgcagctt actcataaca agatcacaaa gctgggctct tttgaaggat 780
tggtaaacct gaccttcatc catctccagc acaatcggct gaaagaggat gctgtttcag 840
ctgcttttaa aggtcttaaa tcactcgaat accttgactt gagcttcaat cagatagcca 900
gactgccttc tggtctccct gtctctcttc taactctcta cttagacaac aataagatca 960
gcaacatccc tratgagtat ttcaagcgtt ttaatgcatt gcagtatctg cgtttatctc1020
acaacgaact ggctgatagt ggaatacctg gaaattcttt caatgtgtca tccctggttg1080
agctggatct gtcctataac aagcttaaaa acataccaac tgtcaatgaa aaccttgaaa1140
actattacct ggaggtcaat caacttgaga agtttgacat aaagagcttc tgcaagatcc1200
tggggccatt atcctactcc aagatcaagc atttgcgttt ggatggcaat cgcatctcag1260
aaaccagtct tecaceggat atgtatgaat gtetaegtgt tgetaaegaa gteaetetta1320
attaatatot gratootgga acaatattit atggttatgt tittototgtgt groagttito1380
atagtateca tattttatta etgtttatta etteeatgaa ttttaaaate tgagggaaat1440
gttttgtaaa catttatttt tttttaaagg aaaaggatgg aaaggccagg gcctaatttc1500
catccaccaa ggaacacacc acattattcc acggaatagg ccatcggg
```

- (2) INFORMATION ON SEQ ID NO. 20:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 844 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: Partial cDNAs produced from individual
 - ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:

acctgcagag gggtccatac ggcgttgttc tgggattcccg tcgtaactta aagggaaatt 60 ttcacaatgt ccggagcct tgatgtcctg caaatgaagg aggaggatgt ccttaagttc120 cttgcagcag gaacccactt aggtggcacc aatcttgact tccagatgga acagtacatc180 tataaaagga aaagtgatgg catctatatc ataaatctca agaggacctg ggagaagctt240 ctgctggcag ctcgtgcaat tgttgccatt gaaaaccctg ctgatgtcag tgttatatcc300 tccaggaata ctggccagag ggctgtgctg aagtttgctg ctgccactgg agccactcca360 attgctggcc gcttcactcc tggaaccttc actaaccaga tccaggcagc cttccgggag420 ccacggcttc ttgtgggttac tgaccccagg gctgaccacc agcctctcac ggaggcatct480 tatgttaacc tacctaccat tgcgctgtgt aacacagatt ctccttcgcg ctatgtggac540 attgcaatcc catgcaacaa caaggtaatg attttaggat ctagagtttg tgaatgcgt600 ctctagaaaa aacattcctg tgcacattgt tagagcttgg agttgaggct actgactggc660 cgatgaactc gcaagtgtag gtagtgtgct acatgagggg caagttttcg ctaacacac720 aagggtctct gccaatga gtggagtttg atagtaattc ttgctacaag tataacatta780 ctgcatgaca gctttgtga gaaatgaaa catttggaaa atagtgtgt ctctgccttg840 tcca

- (2) INFORMATION ON SEQ ID NO. 21:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 862 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

totatgocat toacgoogag gagatootgg agaagcacco gogagggggo agottoggat660 actgottog cotggootgg gtggoottoo coctogocot ggtcagoggo atcatotaca720 tocacctacg gaagcgggag tgagcgooco gootogotog gotgoococg cocottocog780 goococotog cogogotoo tocaaaaaaa aaaccttaa cogoggggaa aaaaaaaaa840 aaaaaaggaag gaaaaaaaaa aa

(2) INFORMATION ON SEQ ID NO. 22:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 546 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) ANTI-SENSE: NO

(iii) HYPOTHETICAL: NO

- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:

(vii) OTHER ORIGIN:

- (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

cccaqccaaq ggtccttcaq gtagqaggtc ctqgqtqact ttqqaaqtcc qtagtqtctc 60 attqcaqata atttttaqct tagqqqctq tqqctaqqtc qgttctctcc tttccaqtcq120 gaqacctctq ccqcaaacat qctccqcaq atcatcqqtc agqccaaqaa qcatccqaqc180 ttqatccccc tctttqtatt tattqqaact qqaqctactq qaqcaacact qtatctcttq240 cqtctqqcat tqttcaatcc aqatqtttqt tqqqacaqaa ataacccaqa qccctqqaac300 aaactqqqtc ccaatqatca atacaaqttc tactcaqtqa atqtqqatta caqcaaqctq360 aqaaqqaac qtccaqattt ccaaatqaaa tqtttcacta taacqctqct ttaqaaatqaa420 qqtcttccaq aagccacatc cqcacaattt tccacttaac caqqaaatat ttctcctctt480 aaatqaatqa aatcaatqqt qqqqqqcc attqqaaqcc ctattqqqt tcaaqtqttq540 aataaa

(2) INFORMATION ON SEQ ID NO. 23:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1591 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual

ESTs by assembling and editing

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:

```
geogaggage egageoegee acceecege eegeeegeeg eegeeatggg etgeeteggg
 aacagtaaga ccgaggacca gcgcaacgag gagaaggcgc aggtgaggcc aacaaaaaga 120
 togagaagca gotgoagaag gacaagcagg totacogggo cacgcaccgo otgotgotgo 180
 tgggtgctgg agaatctggt aaaagcacca ttgtgaagca gatgaggatc ctgcatgtta 240
 atgggtttaa tggagacagt gagaaggcaa ccaaagtgca ggacatcaaa aacaacctga 300
 aagaggcgat tgaaaccatt gtggccgcca tgagcaacct ggtgcccccc gtggagctgg 360
 ccaaccccga gaaccagttc agagtggact acattctgag tgtgatgaac gtgcctgact 420
 ttgacttccc tcccgaattc tatgagcatg ccaaggctct gtgggaggat gaaggagtgc 480
gtgcctgcta cgaacgctcc aacgagtacc agctgattga ctgtgcccag tacttcctgg 540
acaagatcga cgtgatcaag caggctgact atgtgccgag cgatcaggac ctgcttcgct 600
googtgtoot gacttotgga atotttgaga coaagttoca ggtggacaaa gtcaacttoo 660
acatgtttga cgtgggtggc cagcgcgatg aacgccgcaa gtggatccag tgcttcaacg 720
atgtgactgc catcatcttc gtggtggcca gcagcagcta caacatggtc atccgggagg 780
acaaccagac caaccgcctg caggaggctc tgaacctctt caagagcatc tggaacaaca 840
gatgqctgcg caccatctct gtgatcctgt tcctcaacaa gcaagatctg ctcgctgaga 900
aagtccttgc tgggaaatcg aagattgagg actactttcc agaatttgct cgctacacta 960
cteetgagga tgetaeteee gageeeggag aggaeeeaeg egtgaeeegg geeaagtaet1020
tcattcgaga tgagtttctg aggatcagca ctgccagtgg agatgggcgt cactactgct1080
acceteattt cacetgeget giggacactg agaacateeg eegigtgite aacgaetgee1140
gtgacatcat tcagcgcatg caccttcgtc agtacgagct gctctaagaa gggaaccccc1200
adatttaatt aaagcottaa goacaattaa ttaaaagtga aacgtaattg tacaagcagt1260
taatcaccca ccatagggca tgattaacaa agcaaccttt cccttccccc gagtgatttt1320
gegaaaeeee ettteeeett eagettgett agatgtteea aatttagaaa gettaaggeg1380
gcctacagaa aaaggaaaaa aggccacaaa agttccctct cactttcagt aaaaataaat1440
aaaacagcag cagcaaacaa ataaaatgaa ataaaagaaa caaatgaaat aaatattgtg1500
ggcaaaaggg gaaagaagaa aaggggggg g
                                                                1591
```

(2) INFORMATION ON SEQ ID NO. 24:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 441 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

ggcaggcaga tacgttcgtc agcttgctcc tttctgcccg tggacgccgc cgaagaagca 60 tcgttaaagt ctctctcac cctgccgtca tgtctaagtc agagtctcct aaagagcccg120 aacaagctgag gaagctcttc attggagggt tgagctttga aacaactgat gagagcctga180 ggagccattt tgagcaatgg ggaacgctca cggactgtgt ggtaatgaga gatccaaaca240 ccaagcgctc caggggcttt gggtttgtca catatgccac tgtggaggag gtggatgcag300 ctatgaatgc aaggccacac aaggtggatg gaagagttgt ggaaccaaaa agagctgttt360 cagagaagat ttgaaaagcc aggtgccact tacctgtgaa aaggtatttg ttggtggatt420 aaggagcact tgagacatca c

(2) INFORMATION ON SEQ ID NO. 25:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1131 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:

(vii) OTHER ORIGIN:

(A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:

```
cgggaggtga aatccggttc taaccggtcc ggggctcca gcgctataaa aactttataa 60 acccccgga gcccgagcag tgtgaagaag aggcgagaac gacccccgga ccgaccaaag 120 cccgcgcgcc gctgcatcc gcgtccagca cctacgtccc gctgccgtcg ccgccgcac 180 catgcccaag agaaaggctg aagggagatgc taaagggagat aaagcaaaagg tgaaaggacga 240 accacagaga agatccgcga ggttgtctgc taaacctgct cctccaaagc cagagcccaa 300 gcctaaaaag gcccctgcaa agaagggaga gaaggtaccc aaagggaaaa agggaaaagc 360 tgatgctgc aaggaggga ataaccctgc agaaaatgga gatgccaaaa cagaccaggc 420 acagaaagct gaaggtgct gagatgccaa gtgaagtgtg tgcatttttg ataactgtgt 480 acttctggtg actgtacagt ttgaaatact atttttatc aagttttata aaaatgcaga 540 attttgtttt tttttttataa agctatgttg ttagcacaca gaacacttca 600 ttgttgtttt tggagaaaa acaccttcc cttctagtt tgaagagact ccccagagga 720
```

gggattccct gactttgaca cacatggcca ccttggcaca aaagccttgt ggtatagaaa 780 aacaaatttg titttatgtc ctcttccc tttccatctt tcagcataga cttaactccc 840 ttaagcccag acatctgttg agacctgacc cctagtcatt ggttaccagt gtgtcaggca 900 atctggactt tccagtgatg ccactgagat ggcacctgtc aaaagagcag tggttccatt 960 tctagattgt ggatcttcag ataaattctg ccattttcat ttcacttcct gaaaagtcaggl020 gtcggcttgt gaaaagttgt taaacaacat gctaaatgtg aaatgtcaac cctcactctal080 aaacttttcc ctgggtcaga ggatccgatg gaggacttca attgggggtt t 1131

(2) INFORMATION ON SEQ ID NO. 26:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1071 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:

```
gtaccctcaa agacagagac accaagaaga atcggaacat acaggctttg atatcaaagg
tttataaagc caatatctgg gaaagagaaa accgtgagac ttccagatct tctctggtga 120
agtgttgttt cttgcaacga tcacgaacat gaacatcaaa ggatcgccat ggaaagggtc 180
cottottgetg etgetggtgt caaacetget cetgtgecag agegtggece cettgeceat 240
ctgtcccggc ggggctgccc gatgccaggt gacccttcga gacctgtttg accgcgccgt 300
egtectgtee eactacatee ataacetete etcagaaatg tteagegaat tegataaaeg 360
gtatacceat ggccgggggt teattaceaa ggccateaac agetgeeaca ettetteeet 420
tgccacccc gaagacaagg agcaagccca acagatgaat caaaaagact ttctgagcct 480
gatagtcage atattgcgat cetggaatga geetetgtat catetggtca eggaagtaeg 540
tggtatgcaa gaageceegg aggetateet atecaaaget gtagagattg aggageaaac 600
caaacqqctt ctagaqqqca tqqaqctqat aqtcaqccaq qttcatcctq aaaccaaaqa 660
aaatgagate taccetgtet ggtegggaet tecatecetg cagatggetg atgaagagte 720
tegeetttet gettattata acetgeteea etgeetaege agggatteae ataaaatega 780
caattatete aageteetga agtgeegaat catecacaac aacaactget aageecacat 840
ccatttcatc tatttctgag aaggtcctta atgatccgtt ccattgcaag cttcttttag 900
ttgtatctct tttgaatcca tgcttgggtg taacaggtct cctcttaaaa aataaaaact 960
gacteettag acateaaa atetaaaaaa aettaatggg eegggegeag tggeteatgg1020
ctgtggtccc ggcactttgg gaggccgagg caggcggatc aggaggtcag g
```

- (2) INFORMATION ON SEQ ID NO. 27:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 896 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

- (2) INFORMATION ON SEQ ID NO. 28:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 1050 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

ttttcatttt ttttttt ttttctcag ttcaagttta atacaaacta caaaagatta 60 atgggttgct ctactaatac atcatacaaa ccagtagcct gcccacaacg ccaactcagg 120 ccatteetac caaaggaaga aaggetggte tetecacee etgtaggaaa ggeetgeett 180 gtaagacacc acaattcggc tgaatctgaa gtcttgtgtt ttactaatgg aaaaaaaaa 240 tacagaagag gttttgttct catggctgcc caccgcagcc tggcactaaa acagcccagc 300 geteaettet gettggagaa atattetttg etettttgga eateaggett gatggtatea 360 ctgccaggtt tccagccage tgggcacaet tccccatgtt tgtcagtgaa ctggaaggcc 420 tgaactagtc tcaaagtctc atccacagag cggccaacág ggaggtcatt tacagtgatc 480 tgccgaagaa taccettate atcaatgata aaaaggcccc tgaacgagat gccttcatca 540 geetttaaga eeccataate etgageaatg gtgegetteg ggtetgatae caaaggaatg 600 ttcatgggtc ccagtcctcc ttgtttctta ggtgtattga cccatgctag atgacagaag 660 tgagaateca cagaageace aateaettgg cagttgagtt tettaaatte ttetgeeeta 720 tractgaaag caatgatote egtggggcar acaaaggtga agtcaagagg gtaaaagaag 780 aacacaacat atttteettt gtagteagae aggetgatat etttaaaetg accatetgge 840 ataacagetg tggetttgaa gttgggggca gggtgeecaa ttttageatt teetgaagae 900 atottoctat cagcagtoco aacacaagto goaqaaacta accacogaca ccaggcaaga 960 acaagacgcg caagagctct ccggggcgct gcctttatag ccagtaggga tctcgccaca1020 gtcggaacgg acgggggtgc cggagtagga 1050

- (2) INFORMATION ON SEQ ID NO. 29:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 581 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual

ESTs by assembling and editing

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

caggetteet tetggeaaca ggegtgggte aegetetege teggtette tgeegeeate 60 ttggtteege gtteeetga caaaatgee ggegaacace agaaacegte cetgetacag120 ageaggagtt geegeagee caggetgaga cagggtetgg aacaagaatet gacagtgatg180 aateagtace agagettgaa gaacaggatt ceaeceagge aaceacacaa caageecage240 tggeggeage agetgaaate gatgaagaac cagteagtaa ageaaaacag agteggagtg300 aaaaagaagge aeggaagget atgteeaaac tgggtetteg geaggttaca ggagttacta360

gagtcactat ccggaaatct aagaatatcc tctttgtcat cacaaaacca gttgtctaca420 agagccctgc ttcagatacg tacatagttt ttggggaagc cagatcgaag attatcccag480 caagcacaac tagcagctgc tgagaagtca agttcaggtg aactgtctca acgttcagga540 aacccccggc ttccactgta gaggggagt aaggggaggg t

- (2) INFORMATION ON SEQ ID NO. 30:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 264 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN

- (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

gggactatgt tgtgagcctg cgaaagaagt ttgtgtggg actgtgggca gtgaatgcgt 60 tgggaacaat atggaaaact gggagctgcc ctcagtttct ccccaagttg gactcacttt120 cggggtgtcc caaaagcctg attccagggc ctgctagccc gaccccggtg acgcctcac180 ccgcgcctgg ccccagcctt cacccgcagt cgccgcctc cggggcacac cctccgccag240 aaaacagccg gcgggcggcg agac 264

- (2) INFORMATION ON SEQ ID NO. 31:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 111 base pairs
 - (B) TYPE: Nucleic acid
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing
 - (iii) HYPOTHETICAL: NO
 - (iii) ANTI-SENSE: NO
 - (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
 - (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:

cggcgaatca cttataaatg gcgccgaagc aggagcccga aggctaaatt gcaggagggg 60
-tgagcgaatg crgtgctttc atgggcctct tacgttgatg aggcaaagta t 111

- (2) INFORMATION ON SEQ ID NO. 32:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 76 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF

- (iii) HYPOTHETICAL: yes
- (vi) ORIGIN
 - (A) ORGANISM: HUMAN
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 32:

PFCEETKTER LWPRCRPPAA VGFSTQNPGV GDSESNLFSL PFLGSKANPI PTHWSSALIF60 NLPSPPFQNT HIPFQN 76

- (2) INFORMATION ON SEQ ID NO. 33:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 72 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33:

SSFLFSFQTQ FHKNRKDKVF SSRQAKPFPH HQSILKIHEE VERSVSGRLK GSSSSNPTAA60 EKIEIEILKI TS

- (2) INFORMATION ON SEQ ID NO. 34:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 70 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 34:

KKLDYFCAEI KNSHCKTKIK IAQIRKPGGA KCQVSKVHFF SLSKRSSTKT ARIKFSVADK60 QSFFHIINQS

- (2) INFORMATION ON SEQ ID NO. 35:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 60 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 35:

SSGPAPGCSP FAGTRKNFPS MVVLERTFLK INYIFLCIPM EFQFIRCSPW PPQNTEVIPA60

- (2) INFORMATION ON SEQ ID NO. 36:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 63 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 36:

ASGVHTETHR YNLLSAKSRK KGWGYLGWLG FDFLLVCLFC TKTVLSFEYR RDISIYMLSN60 QDG 63

- (2) INFORMATION ON SEQ ID NO. 37:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 170 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes

- (vi) ORIGIN
 - (A) ORGANISM: HUMAN
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 37:

ARAARAAQTP HLTLPADLQT LHLNRPTLSP ESKLEWNNDI PEVNHLNSEH WRKTEKWTGH 60 EETNHLETDF SGDGMTELEL GPSPRLQPIR RHPKELPQYG GPGKDIFEDQ LYLPVHSDGI120 SVHQMFTMAT AEHRSNSSIA GKMLTKVEKN HEKEKSQHLE GSASSSLSSD 170

- (2) INFORMATION ON SEQ ID NO. 38:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 144 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN.
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 38:

ARAPTLDMRF RRRLSADPHA TQRNSAEARG TMDGRVQLMK ALLAGPLRPA ARRWRNPIPF 60 PETFDGDTDR LPEFIVQTCS YMFVDENTFS NDALKVTFLI TRLTGPALQW VIPYIRKESP120 LLNDYRGFLA EMKRVFGWEE DEDF

- (2) INFORMATION ON SEQ ID NO. 39:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 178 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 39:

HSLGRAPVET LAVATGTANS SQSTRPQARG SPGLEVLVLL PSKDSLHLGQ KAPVIIEQGA 60 LLPDVGDHPL QGWPREAGDE ERHLQGVVGE RVLVHEHVGA RLHDELRESV GISVKRLGKG120 NRVPPATRRG PEGPGQEGLH QLHPTVHRAA RLRGVSLGCV GVSAKASPEA HVEGGGPG 178

- (2) INFORMATION ON SEQ ID NO. 40:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 89 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40:

KLTGINTGCR NMLALCIRGH AQQIQEIYLA TFSRKGTLGI IHYILEVFLG FFFFFLRQSC60 CIAQAGSVVA QSQLIASSIT QGLSNPPTL 89

- (2) INFORMATION ON SEQ ID NO. 41:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 95 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41:

IVTWRKVPMS LCQRPPPFVR IGIFRLLKGL AHIRCDLFIP VVMEGHICQS LESAKAGTRF60 PGPQWGCANP RELGCKFVKN QHHVWQLSIG ARSLP 95

- (2) INFORMATION ON SEQ ID NO. 42:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 154 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF

(iii) HYPOTHETICAL: yes

(vi) ORIGIN

(A) ORGANISM: HUMAN

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 42:

CQLVFRIQTD GSYWSLGLTS SGNITFSWAE MLLPALKQHS VLKTSWQAPG SNTQLPNMML 60 ILHEFATQFS RVCTPPLWAG EPGPGLRRLQ ALADVALHNN GNEKVTPYVR QALKESEYPN120 PHKRRGTLAK THGNFPPSND LDRRATQDSP SCSV 154

(2) INFORMATION ON SEQ ID NO. 43:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 79 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: ORF
- (iii) HYPOTHETICAL: yes
- (vi) ORIGIN
 - (A) ORGANISM: HUMAN
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 43:

LASTLGVETC LPYVSEDMLS RSKRYIWQLF LEKAHWVSFI TFLSFFGFFF FFFETVLLYC60 PGWSVVAQSQ LIASSITQA 79

- (2) INFORMATION ON SEQ ID NO. 44:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 82 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 44:

CQLVFRIQTD GSYWSLGLTS SGNITFSWAE MLLPALKQHS VLKTSWQAPG SNTQLPNMML60 ILHEFATSWL PRLQHSAVGT QS 82

- (2) INFORMATION ON SEQ ID NO. 45:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 68 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 45:

RGSKDRNSGQ GSGSYGQLSC RGFSDQFSRV CTPPLWAGEP GPGLRRLQAL ADVALHNNGN60 EKVTPYVR 68

- (2) INFORMATION ON SEQ ID NO. 46:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 87 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 46:

DYVVSLRKKF VWGLWAVNAL GTIWKTGSCP QFLPKLOSLS GCPKSLIPGP ASPTPVTPPP60 APGPSLHPRS PSGAHPPPE NSRRAAR 87

- (2) INFORMATION ON SEQ ID NO. 47:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 51 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes

(vi) ORIGIN (A) ORGANISM: HUMAN	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 47:	
QALESGFWDT PKVSPTWGET EGSSQFSILF PTHSLPTVPT QTSFAGSQHS P	51
(2) INFORMATION ON SEQ ID NO. 48:	
 (i) SEQUENCE CHARACTERISTIC: (A) LENGTH: 20 amino acids (B) TYPE: Protein (C) STRAND: individual (D) TOPOLOGY: linear 	•
(ii) MOLECULE TYPE: ORF	
(iii) HYPOTHETICAL: yes	
(vi) ORIGIN (A) ORGANISM: HUMAN	•
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 48:	
RRITYKWRRS RSPKAKLQEG	20
(2) INFORMATION ON SEQ ID NO. 49:	
 (i) SEQUENCE CHARACTERISTIC: (A) LENGTH: 36 amino acids (B) TYPE: Protein (C) STRAND: individual (D) TOPOLOGY: linear 	
(ii) MOLECULE TYPE: ORF	
(iii) HYPOTHETICAL: yes	
(vi) ORIGIN (A) ORGANISM: HUMAN	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 49:

(2) INFORMATION ON SEQ ID NO. 50: (i) SEQUENCE CHARACTERISTIC: (A) LENGTH: 26 amino acids (B) TYPE: Protein (C) STRAND: individual (D) TOPOLOGY: linear (ii) MOLECULE TYPE: ORF (iii) HYPOTHETICAL: yes (vi) ORIGIN (A) ORGANISM: HUMAN (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 50: 26 KHSIRSPLLQ FSLRAPASAP FISDSP (2) INFORMATION ON SEQ ID NO. 51: (i) SEQUENCE CHARACTERISTIC: (A) LENGTH: 25 amino acids (B) TYPE: Protein (C) STRAND: individual (D) TOPOLOGY: linear (ii) MOLECULE TYPE: ORF (iii) HYPOTHETICAL: yes (vi) ORIGIN

EAHESTAFAH PSCNLAFGLL LRRHL

25

- (2) INFORMATION ON SEQ ID NO. 52:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 3665 base pairs

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 51:

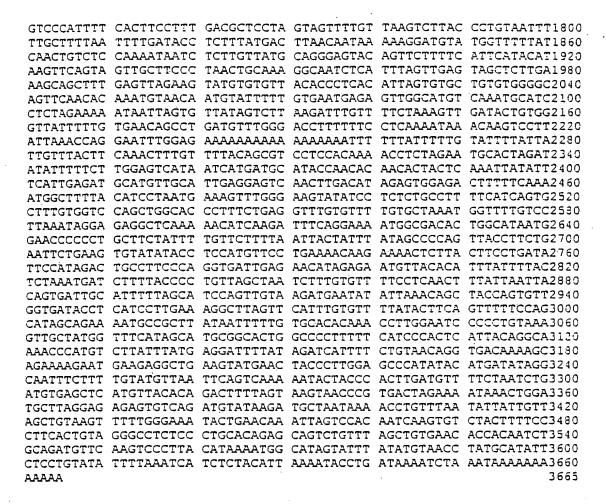
- (B) TYPE: Nucleic acid
- (C) STRAND: individual
- (D) TOPOLOGY: linear

(A) ORGANISM: HUMAN

(ii) MOLECULE TYPE: Partial cDNAs produced from individual ESTs by assembling and editing

- (iii) HYPOTHETICAL: NO
- (iii) ANTI-SENSE: NO
- (vi) ORIGIN:
 - (A) ORGANISM: HUMAN
 - (C) ORGAN:
- (vii) OTHER ORIGIN:
 - (A) LIBRARY: cDNA library
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 52:

	TGAAGAGACG	NACACTGAGC	GGTTGTGGCC	GCGTTGCCGA	CCTCCAGCAG	60
GGCCATTTTG	TGAAGAGACG	, =		AGAAATCGAA		L20
CAGTCGGCTT	CTCTACGCAG			ACTGGAGCTC		180
CTCCCCTTCT	TGGGCAGCAA	CTTCCAGAAC	ACACACATTC	CATTCCAAAA	CTGATTTTAT	240
TTTAACCTCC	CTTCCCCACC	GATGCAACTT	GGTGTGCACT	ACAGCAAATG	TACAGGTGTT	300
AAAGACATTT	TAAACATAAT		TGGATTTAAG	ATGTAATTTT	TAAAATTTCT	360
AATTTTTTTA	TIGTTTCCAA		GAGGAGGAGG	AGCCTTTTAG	CCTCTCATAA	420
ATTTCTATTT	TTTCTGCAGC		AAGATTGATT	GATGATGTGG	AAAGGGCTTT	480
ACTGACCTCT	CTACTTCCTC	GTGTATTTTT	GGTTTTTGTG	GAAACTGCTT	TTGGAAAGAG	540
GCTTGTCTGC	TACTGAAAAC	TTTATCCTGC	TTTGCACCTC	CCGTTTTTCT	AATCTGGGCT	600
AAAAGAAATG	AACTTTACTG	ACTTGACATT		TTCAGCTTAC		660
ATTTTTATTT	TTGTTTTTT	ACAGTGAGAT	TTTTTTGATC	AACGTTACCA		720
TTTGTGAGGA	AACCTTTACC	CATCAAACAC		AACACTCTTG		780
TCCTCGCTCC	ATGAACTCCC	GTGTATTCAT	TGGGAATCTC	GTGGGCTGCT		840
ATCTGATGTG	GAGGCAATCT	TTTCGAAGTA	TGGCAAAATT	CGGGCTGCTG	TAGCAGGAGA	900
GGGCTTTGCC	TTCGTTCAGT	ATGTTAATGA	GAGAAATGCC	CTGGCTGCAG		960
	ATGATTGCTG	GCCAGGTTTT	AGATATTAAC			020
GAACCGAGGA	AAAGCAGGTG	TGAAACGATC	TGCAGCGGAG			
CTTGGACTAT	GACTTTCAAC	GGGACTATTA	TGATAGGATG			
ACCTCCTCCT	CCTCCTATTG		AGTGCCCTCG			
	CGAAGGGGCA	AAAGTGGCTT	CAATTCTAAG			
CAAGTCTGGA	AAGTTGAAAG	GAGATGACCT	TCAGGCCATT			
	GTGGATTCTC	TCCTGGAAAA		ATTGAAAAGG		
ACAAGCAGTA					GCTCCGTGAA	1440
GAAAGATGAG			TGAGGGGGGT			
GGGGGACCTA			AGATCGGGGG			
CAAGGATGAT			AGAGGATGAC			
GGATGACTCT		GTGGGGTTTA				
GGAIGACICI GCGCTTGTCI				CTAGTATCT	r cagcacatgc	
TCACTGTTCT		GTCCTTCCC	TGTTCATTAR	TTCATATTG	CCCGCGCCTA	1 / 4 U
ICACIGITO	COCCARGO				•	

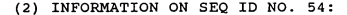


(2) INFORMATION ON SEQ ID NO. 53:

- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 301 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: ORF
- (iii) HYPOTHETICAL: yes
- (vi) ORIGIN
 - (A) ORGANISM: HUMAN
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 53:

GNLYPSNTMA SNVTNKTDPR SMNSRVFIGN LNTLVVKKSD VEAIFSKYGK IVGCSVHKGF 60 AFVQYVNERN ARAAVAGEDG RMIAGQVLDI NLAAEPKVNR GKAGVKRSAA EMYGSSFDLD120 YDFQRDYYDR MYSYPARVPP PPPIARAVVP SKRQRVSGNT SRRGKSGFNS KSGQRGSSKS180 GKLKGDDLQA IKKELTQIKQ KVDSLLENLE KIEKEQSKQA VEMKNDKSEE EQSSSSVKKD240 ETNVKMESEG GADDSAEEGD LLDDDDNEDR GDDQLELIKD DEKEAEEGED DRDKANGEDD300

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- (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 112 amino acids

 - (B) TYPE: Protein(C) STRAND: individual
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: ORF
- (iii) HYPOTHETICAL: yes
- (vi) ORIGIN
 - (A) ORGANISM: HUMAN
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 54:

ESSSPLALSL SSSPSSASFS SSLINSSWSS PRSSLSSSSS RSPSSAESSA PPSDSIFTLV 60 SSFFTELLLC SSSDLSFFIS TACLLCSFSI FSRFSRREST FCFIWVSSFL MA

- (2) INFORMATION ON SEQ ID NO. 55:
 - (i) SEQUENCE CHARACTERISTIC:
 - (A) LENGTH: 107 amino acids
 - (B) TYPE: Protein
 - (C) STRAND: individual
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: ORF
 - (iii) HYPOTHETICAL: yes
 - (vi) ORIGIN
 - (A) ORGANISM: HUMAN
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 55: